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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Comprehensive Assessment Information Rule

REPORTING FORM

When completed, send this form to:

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Office of Toxic Substances, TS-790
U.S. Environmental Protection Agency
401 M Street, SW
Washington, DC 20460
Attention: CAIR Reporting Office

For Agency Use Only:

Date of Receipt: _____

Document
Control Number:

Docket Number:

SECTION 1 GENERAL MANUFACTURER, IMPORTER, AND PROCESSOR INFORMATION

PART A GENERAL REPORTING INFORMATION

1.01 This Comprehensive Assessment Information Rule (CAIR) Reporting Form has been

completed in response to the Federal Register Notice of..... [1][2] [2][2] [8][8]
mo. day year

CBI

☐ a. If a Chemical Abstracts Service Number (CAS No.) is provided in the Federal Register, list the CAS No. [0][2][6][4][7][1]-[6][2]-[5]

b. If a chemical substance CAS No. is not provided in the Federal Register, list either (i) the chemical name, (ii) the mixture name, or (iii) the trade name of the chemical substance as provided in the Federal Register.

(i) Chemical name as listed in the rule NA

(ii) Name of mixture as listed in the rule

(iii) Trade name as listed in the rule

c. If a chemical category is provided in the Federal Register, report the name of the category as listed in the rule, the chemical substance CAS No. you are reporting on which falls under the listed category, and the chemical name of the substance you are reporting on which falls under the listed category.

Name of category as listed in the rule NA

CAS No. of chemical substance [][][][][][]-[][]-[]

Name of chemical substance

1.02 Identify your reporting status under CAIR by circling the appropriate response(s).

CBI Manufacturer 1

☐ Importer 2

Processor 3

X/P manufacturer reporting for customer who is a processor 4

X/P processor reporting for customer who is a processor 5

☐ Mark (X) this box if you attach a continuation sheet.

1.03 Does the substance you are reporting on have an "x/p" designation associated with it in the above-listed Federal Register Notice?

CBI
☐ Yes [X] Go to question 1.04
☐ No [☐] Go to question 1.05

1.04 a. Do you manufacture, import, or process the listed substance and distribute it under a trade name(s) different than that listed in the Federal Register Notice? Circle the appropriate response.

CBI
☐ Yes 1
☐ No (2)

b. Check the appropriate box below: NA

☐ You have chosen to notify your customers of their reporting obligations
Provide the trade name(s)

☐ You have chosen to report for your customers

☐ You have submitted the trade name(s) to EPA one day after the effective date of the rule in the Federal Register Notice under which you are reporting.

1.05 If you buy a trade name product and are reporting because you were notified of your reporting requirements by your trade name supplier, provide that trade name.

CBI
Trade name LUPRANATE T80 MONDUR TD, VORANATE T-80, RUBINATE TDI

☐ Is the trade name product a mixture? Circle the appropriate response.

Yes 1
No (2)

1.06 Certification -- The person who is responsible for the completion of this form must sign the certification statement below:

CBI
☐ "I hereby certify that, to the best of my knowledge and belief, all information entered on this form is complete and accurate."

PAUL MCINTYRE
NAME


SIGNATURE

6/22/89
DATE SIGNED

PLANT MANAGER
TITLE

(717) 542 - 4171
TELEPHONE NO.

☐ Mark (X) this box if you attach a continuation sheet.

- 1.07 Exemptions From Reporting -- If you have provided EPA or another Federal agency with the required information on a CAIR Reporting Form for the listed substance within the past 3 years, and this information is current, accurate, and complete for the time period specified in the rule, then sign the certification below. You CBI ☐ are required to complete section 1 of this CAIR form and provide any information now required but not previously submitted. Provide a copy of any previous submissions along with your Section 1 submission.

"I hereby certify that, to the best of my knowledge and belief, all required information which I have not included in this CAIR Reporting Form has been submitted to EPA within the past 3 years and is current, accurate, and complete for the time period specified in the rule."

_____ NA NAME	_____ SIGNATURE	_____ DATE SIGNED
_____ TITLE	(_____) - TELEPHONE NO.	_____ DATE OF PREVIOUS SUBMISSION

- 1.08 CBI Certification -- If you have asserted any CBI claims in this report you must certify that the following statements truthfully and accurately apply to all of those confidentiality claims which you have asserted.

CBI ☐ "My company has taken measures to protect the confidentiality of the information, and it will continue to take these measures; the information is not, and has not been, reasonably ascertainable by other persons (other than government bodies) by using legitimate means (other than discovery based on a showing of special need in a judicial or quasi-judicial proceeding) without my company's consent; the information is not publicly available elsewhere; and disclosure of the information would cause substantial harm to my company's competitive position."

_____ NA NAME	_____ SIGNATURE	_____ DATE SIGNED
_____ TITLE	(_____) - TELEPHONE NO.	

☐ Mark (X) this box if you attach a continuation sheet.

1.09 Facility Identification

1.10 Company Headquarters Identification

☐ Mark (X) this box if you attach a continuation sheet.

1.11 Parent Company Identification

CBI Name [P][M][C], [I][N][C] _____
[] Address [P][O] [B][O]X [1][3][6][7] _____
 Street
 [S][U][N] [V][A][L][L][E][Y] _____
 City
 [C][A] [9][1][3][5][2]--[][][]
 State Zip

Dun & Bradstreet Number [0][7]-[6][1][9]-[1][5][1][9]

1.12 Technical Contact

[illegible]

1.13 This reporting year is from

0	1
Mo.	Year

 to

1	2
Mo.	Year

☐ Mark (X) this box if you attach a continuation sheet.

[illegible]

[][] [][][][]--[][][][]
State Zip

Date of Sale [N] [A] [] [] [] []
Mo. Day Year

Telephone Number[][]-[][]-[][]

[illegible]

--
State Zip

Date of Purchase [] [] [] [] [] []
Mo. Day Year

Telephone Number () - () - ()

8

1.16 For each classification listed below, state the quantity of the listed substance that was manufactured, imported, or processed at your facility during the reporting year.

CBI

<u>Classification</u>	<u>Quantity (kg/yr)</u>
<input type="checkbox"/> Manufactured	NA
Imported	NA
Processed (include quantity repackaged)	254,000
Of that quantity manufactured or imported, report that quantity:	
In storage at the beginning of the reporting year	NA
For on-site use or processing	NA
For direct commercial distribution (including export)	NA
In storage at the end of the reporting year	NA
Of that quantity processed, report that quantity:	
In storage at the beginning of the reporting year	14,000
Processed as a reactant (chemical producer)	NA
Processed as a formulation component (mixture producer)	NA
Processed as an article component (article producer)	264,000
Repackaged (including export)	NA
In storage at the end of the reporting year	5,000

☐ Mark (X) this box if you attach a continuation sheet.

1.17 Mixture -- If the listed substance on which you are required to report is a mixture or a component of a mixture, provide the following information for each component chemical. (If the mixture composition is variable, report an average percentage of each component chemical for all formulations.)

[]

☐ Mark (X) this box if you attach a continuation sheet.

2.04 State the quantity of the listed substance that your facility manufactured, imported, or processed during the 3 corporate fiscal years preceding the reporting year in descending order.

CBI

☐ Year ending [1][2] [8][7]
Mo. Year

Quantity manufactured NA kg

Quantity imported NA kg

Quantity processed 256,000 kg

Year ending [1][2] [8][6]
Mo. Year

Quantity manufactured NA kg

Quantity imported NA kg

Quantity processed 128,000 kg

Year ending [1][2] [8][5]
Mo. Year

Quantity manufactured NA kg

Quantity imported NA kg

Quantity processed NA kg

2.05 Specify the manner in which you manufactured the listed substance. Circle all appropriate process types.

CBI

☐ Continuous process NA 1
Semicontinuous process 2
Batch process 3

☐ Mark (X) this box if you attach a continuation sheet.

2.06 Specify the manner in which you processed the listed substance. Circle all appropriate process types.

☐ Continuous process 1
Semicontinuous process 2
Batch process (3)

2.07 State your facility's name-plate capacity for manufacturing or processing the listed substance. (If you are a batch manufacturer or batch processor, do not answer this question.)

☐ Manufacturing capacity NA kg/yr
Processing capacity UK kg/yr

2.08 If you intend to increase or decrease the quantity of the listed substance manufactured, imported, or processed at any time after your current corporate fiscal year, estimate the increase or decrease based upon the reporting year's production volume.

<input type="checkbox"/>	Manufacturing Quantity (kg)	Importing Quantity (kg)	Processing Quantity (kg)
Amount of increase	NA	NA	NA
Amount of decrease	NA	NA	NA

☐ Mark (X) this box if you attach a continuation sheet.

2.09 For the three largest volume manufacturing or processing process types involving the listed substance, specify the number of days you manufactured or processed the listed substance during the reporting year. Also specify the average number of hours per day each process type was operated. (If only one or two operations are involved, list those.)

CBI

☐

	<u>Days/Year</u>	<u>Average Hours/Day</u>
Process Type #1 (The process type involving the largest quantity of the listed substance.)		
Manufactured	<u>NA</u>	<u>NA</u>
Processed	<u>260</u>	<u>22</u>
Process Type #2 (The process type involving the 2nd largest quantity of the listed substance.)		
Manufactured	<u>NA</u>	<u>NA</u>
Processed	<u>NA</u>	<u>NA</u>
Process Type #3 (The process type involving the 3rd largest quantity of the listed substance.)		
Manufactured	<u>NA</u>	<u>NA</u>
Processed	<u>NA</u>	<u>NA</u>

2.10 State the maximum daily inventory and average monthly inventory of the listed substance that was stored on-site during the reporting year in the form of a bulk chemical.

CBI

☐

Maximum daily inventory kg

Average monthly inventory kg

☐ Mark (X) this box if you attach a continuation sheet.

2.11 **Related Product Types** -- List any byproducts, coproducts, or impurities present with the listed substance in concentrations greater than 0.1 percent as it is manufactured, imported, or processed. The source of byproducts, coproducts, or impurities means the source from which the byproducts, coproducts, or impurities are made or introduced into the product (e.g., carryover from raw material, reaction product, etc.).

CBI

☐

<u>CAS No.</u>	<u>Chemical Name</u>	<u>Byproduct, Coproduct or Impurity¹</u>	<u>Concentration (%) (specify \pm % precision)</u>	<u>Source of By-products, Coproducts, or Impurities</u>
NA				

¹Use the following codes to designate byproduct, coproduct, or impurity:

B = Byproduct
C = Coproduct
I = Impurity

☐ Mark (X) this box if you attach a continuation sheet.

- 2.12 Existing Product Types -- List all existing product types which you manufactured, imported, or processed using the listed substance during the reporting year. List the quantity of listed substance you use for each product type as a percentage of the total volume of listed substance used during the reporting year. Also list the quantity of listed substance used captively on-site as a percentage of the value listed under column b., and the types of end-users for each product type. (Refer to CBI ☐ the instructions for further explanation and an example.)

a. Product Types ¹	b. % of Quantity Manufactured, Imported, or Processed	c. % of Quantity Used Captively On-Site	d. Type of End-Users ²
K	100	100	I

¹Use the following codes to designate product types:

A = Solvent	L = Moldable/Castable/Rubber and additives
B = Synthetic reactant	M = Plasticizer
C = Catalyst/Initiator/Accelerator/ Sensitizer	N = Dye/Pigment/Colorant/Ink and additives
D = Inhibitor/Stabilizer/Scavenger/ Antioxidant	O = Photographic/Reprographic chemical and additives
E = Analytical reagent	P = Electrodeposition/Plating chemicals
F = Chelator/Coagulant/Sequestrant	Q = Fuel and fuel additives
G = Cleanser/Detergent/Degreaser	R = Explosive chemicals and additives
H = Lubricant/Friction modifier/Antiwear agent	S = Fragrance/Flavor chemicals
I = Surfactant/Emulsifier	T = Pollution control chemicals
J = Flame retardant	U = Functional fluids and additives
K = Coating/Binder/Adhesive and additives	V = Metal alloy and additives
	W = Rheological modifier
	X = Other (specify) _____

²Use the following codes to designate the type of end-users:

I = Industrial	CS = Consumer
CM = Commercial	H = Other (specify) _____

☐ Mark (X) this box if you attach a continuation sheet.

- 2.13 Expected Product Types -- Identify all product types which you expect to manufacture, import, or process using the listed substance at any time after your current corporate fiscal year. For each use, specify the quantity you expect to manufacture, import, or process for each use as a percentage of the total volume of listed substance used during the reporting year. Also list the quantity of listed substance used captively on-site as a percentage of the value listed under column b., and the types of end-users for each product type. (Refer to the instructions for further explanation and an example.)

CBI

☐

a.	b.	c.	d.
Product Types ¹	% of Quantity Manufactured, Imported, or Processed	% of Quantity Used Captively On-Site	Type of End-Users ²
K	100	100	I

¹Use the following codes to designate product types:

A = Solvent	L = Moldable/Castable/Rubber and additives
B = Synthetic reactant	M = Plasticizer
C = Catalyst/Initiator/Accelerator/ Sensitizer	N = Dye/Pigment/Colorant/Ink and additives
D = Inhibitor/Stabilizer/Scavenger/ Antioxidant	O = Photographic/Reprographic chemical and additives
E = Analytical reagent	P = Electrodeposition/Plating chemicals
F = Chelator/Coagulant/Sequestrant	Q = Fuel and fuel additives
G = Cleanser/Detergent/Degreaser	R = Explosive chemicals and additives
H = Lubricant/Friction modifier/Antiwear agent	S = Fragrance/Flavor chemicals
I = Surfactant/Emulsifier	T = Pollution control chemicals
J = Flame retardant	U = Functional fluids and additives
K = Coating/Binder/Adhesive and additives	V = Metal alloy and additives
	W = Rheological modifier
	X = Other (specify) _____

²Use the following codes to designate the type of end-users:

I = Industrial	CS = Consumer
CM = Commercial	H = Other (specify) _____

☐ Mark (X) this box if you attach a continuation sheet.

2.14 Final Product -- Complete the following table for each type of final product manufactured, imported, or processed at your facility that contains the listed substance other than as an impurity.

☐

a.	b.	c.	d.
Product Type ¹	Final Product's Physical Form ²	Average % Composition of Listed Substance in Final Product	Type of End-Users ³
NA			

¹Use the following codes to designate product types:

A = Solvent	L = Moldable/Castable/Rubber and additives
B = Synthetic reactant	M = Plasticizer
C = Catalyst/Initiator/Accelerator/Sensitizer	N = Dye/Pigment/Colorant/Ink and additives
D = Inhibitor/Stabilizer/Scavenger/Antioxidant	O = Photographic/Reprographic chemical and additives
E = Analytical reagent	P = Electrodeposition/Plating chemicals
F = Chelator/Coagulant/Sequestrant	Q = Fuel and fuel additives
G = Cleanser/Detergent/Degreaser	R = Explosive chemicals and additives
H = Lubricant/Friction modifier/Antiwear agent	S = Fragrance/Flavor chemicals
I = Surfactant/Emulsifier	T = Pollution control chemicals
J = Flame retardant	U = Functional fluids and additives
K = Coating/Binder/Adhesive and additives	V = Metal alloy and additives
	W = Rheological modifier
	X = Other (specify) _____

²Use the following codes to designate the final product's physical form:

A = Gas	F2 = Crystalline solid
B = Liquid	F3 = Granules
C = Aqueous solution	F4 = Other solid
D = Paste	G = Gel
E = Slurry	H = Other (specify) _____
F1 = Powder	

³Use the following codes to designate the type of end-users:

I = Industrial	CS = Consumer
CM = Commercial	H = Other (specify) _____

☐ Mark (X) this box if you attach a continuation sheet.

2.15 Circle all applicable modes of transportation used to deliver bulk shipments of the
CBI listed substance to off-site customers.

☐ Truck 1
Railcar 2
Barge, Vessel 3
Pipeline 4
Plane 5
Other (specify) NA 6

2.16 Customer Use -- Estimate the quantity of the listed substance used by your customers
CBI or prepared by your customers during the reporting year for use under each category
of end use listed (i-iv).

☐

Category of End Use

i. Industrial Products

Chemical or mixture NA kg/yr

Article NA kg/yr

ii. Commercial Products

Chemical or mixture NA kg/yr

Article NA kg/yr

iii. Consumer Products

Chemical or mixture NA kg/yr

Article NA kg/yr

iv. Other

Distribution (excluding export) NA kg/yr

Export NA kg/yr

Quantity of substance consumed as reactant NA kg/yr

Unknown customer uses NA kg/yr

☐ Mark (X) this box if you attach a continuation sheet.

SECTION 3 PROCESSOR RAW MATERIAL IDENTIFICATION

PART A GENERAL DATA

- 3.01 Specify the quantity purchased and the average price paid for the listed substance for each major source of supply listed. Product trades are treated as purchases.
CBI The average price is the market value of the product that was traded for the listed substance.

☐

<u>Source of Supply</u>	<u>Quantity (kg)</u>	<u>Average Price (\$/kg)</u>
The listed substance was manufactured on-site.	NA	NA
The listed substance was transferred from a different company site.	NA	NA
The listed substance was purchased directly from a manufacturer or importer.	245,000	\$2.00
The listed substance was purchased from a distributor or repackager.	NA	NA
The listed substance was purchased from a mixture producer.	NA	NA

- 3.02 Circle all applicable modes of transportation used to deliver the listed substance to your facility.

- ☐ Truck 1
- Railcar 2
- Barge, Vessel 3
- Pipeline 4
- Plane 5
- Other (specify) _____ 6

☐ Mark (X) this box if you attach a continuation sheet.

3.03 a. Circle all applicable containers used to transport the listed substance to your
CBI facility.

☐ Bags 1
Boxes 2
Free standing tank cylinders 3
Tank rail cars 4
Hopper cars 5
Tank trucks 6
Hopper trucks 7
Drums 8
Pipeline 9
Other (specify) _____ 10

b. If the listed substance is transported in pressurized tank cylinders, tank rail cars, or tank trucks, state the pressure of the tanks.

Tank cylinders NA mmHg
Tank rail cars NA mmHg
Tank trucks 760-7600 mmHg

☐ Mark (X) this box if you attach a continuation sheet.

PART B RAW MATERIAL IN THE FORM OF A MIXTURE

3.04 If you obtain the listed substance in the form of a mixture, list the trade name(s) of the mixture, the name of its supplier(s) or manufacturer(s), an estimate of the average percent composition by weight of the listed substance in the mixture, and the amount of mixture processed during the reporting year.

CBI

☐

<u>Trade Name</u>	<u>Supplier or Manufacturer</u>	<u>Average % Composition by Weight (specify \pm % precision)</u>	<u>Amount Processed (kg/yr)</u>
NA			

☐ Mark (X) this box if you attach a continuation sheet.

PART C RAW MATERIAL VOLUME

3.05 State the quantity of the listed substance used as a raw material during the reporting year in the form of a class I chemical, class II chemical, or polymer, and the percent composition, by weight, of the listed substance.

☐

	Quantity Used (kg/yr)	% Composition by Weight of Listed Sub- stance in Raw Material (specify \pm % precision)
Class I chemical	245,000	99.9%
Class II chemical	NA	NA
Polymer	NA	NA

☐ Mark (X) this box if you attach a continuation sheet.

SECTION 4 PHYSICAL/CHEMICAL PROPERTIES

General Instructions:

If you are reporting on a mixture as defined in the glossary, reply to questions in Section 4 that are inappropriate to mixtures by stating "NA -- mixture."

For questions 4.06-4.15, if you possess any hazard warning statement, label, MSDS, or other notice that addresses the information requested, you may submit a copy or reasonable facsimile in lieu of answering those questions which it addresses.

PART A PHYSICAL/CHEMICAL DATA SUMMARY

- 4.01 Specify the percent purity for the three major¹ technical grade(s) of the listed substance as it is manufactured, imported, or processed. Measure the purity of the substance in the final product form for manufacturing activities, at the time you import the substance, or at the point you begin to process the substance.

☐ CBI

	<u>Manufacture</u>	<u>Import</u>	<u>Process</u>
Technical grade #1	<u>NA</u> % purity	<u>NA</u> % purity	<u>99.9</u> % purity
Technical grade #2	<u>NA</u> % purity	<u>NA</u> % purity	<u>NA</u> % purity
Technical grade #3	<u>NA</u> % purity	<u>NA</u> % purity	<u>NA</u> % purity

¹Major = Greatest quantity of listed substance manufactured, imported or processed.

-
- 4.02 Submit your most recently updated Material Safety Data Sheet (MSDS) for the listed substance, and for every formulation containing the listed substance. If you possess an MSDS that you developed and an MSDS developed by a different source, submit your version. Indicate whether at least one MSDS has been submitted by circling the appropriate response.

Yes (1)

No 2

Indicate whether the MSDS was developed by your company or by a different source.

Your company 1

Another source (2)

☐ Mark (X) this box if you attach a continuation sheet.

4.03 Submit a copy or reasonable facsimile of any hazard information (other than an MSDS) that is provided to your customers/users regarding the listed substance or any formulation containing the listed substance. Indicate whether this information has been submitted by circling the appropriate response.

Yes 1
 No (2)

4.04 For each activity that uses the listed substance, circle all the applicable number(s) corresponding to each physical state of the listed substance during the activity listed. Physical states for importing and processing activities are determined at the time you import or begin to process the listed substance. Physical states for manufacturing, storage, disposal and transport activities are determined using the final state of the product.

CBI
☐

Activity	Physical State				
	Solid	Slurry	Liquid	Liquified Gas	Gas
Manufacture	1	2	3	4	5
Import	1	2	3	4	5
Process	1	2	(3)	4	5
Store	1	2	(3)	4	5
Dispose	1	2	3	4	5
Transport	1	2	3	4	5

☐ Mark (X) this box if you attach a continuation sheet.

4.05 Particle Size -- If the listed substance exists in particulate form during any of the following activities, indicate for each applicable physical state the size and the percentage distribution of the listed substance by activity. Do not include particles ≥ 10 microns in diameter. Measure the physical state and particle sizes for importing and processing activities at the time you import or begin to process the listed substance. Measure the physical state and particle sizes for manufacturing storage, disposal and transport activities using the final state of the product.

CBI

☐

<u>Physical State</u>		<u>Manufacture</u>	<u>Import</u>	<u>Process</u>	<u>Store</u>	<u>Dispose</u>	<u>Transport</u>
Dust	<1 micron	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>
	1 to <5 microns	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>
	5 to <10 microns	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>
Powder	<1 micron	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>
	1 to <5 microns	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>
	5 to <10 microns	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>
Fiber	<1 micron	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>
	1 to <5 microns	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>
	5 to <10 microns	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>
Aerosol	<1 micron	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>
	1 to <5 microns	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>
	5 to <10 microns	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>

☐ Mark (X) this box if you attach a continuation sheet.

SECTION 5 ENVIRONMENTAL FATE

PART A RATE CONSTANTS AND TRANSFORMATION PRODUCTS

5.01 Indicate the rate constants for the following transformation processes.

a. Photolysis:

Absorption spectrum coefficient (peak) UK (1/M cm) at _____ nm
Reaction quantum yield, ϕ UK at _____ nm
Direct photolysis rate constant, k_p , at ... UK 1/hr _____ latitude

b. Oxidation constants at 25°C:

For 1O_2 (singlet oxygen), k_{ox} UK 1/M hr
For RO_2 (peroxy radical), k_{ox} UK 1/M hr

c. Five-day biochemical oxygen demand, BOD_5 ... UK mg/l

d. Biotransformation rate constant:

For bacterial transformation in water, k_b ... UK 1/hr
Specify culture UK

e. Hydrolysis rate constants:

For base-promoted process, k_B UK 1/M hr
For acid-promoted process, k_A UK 1/M hr
For neutral process, k_N UK 1/hr

f. Chemical reduction rate (specify conditions) UK

g. Other (such as spontaneous degradation) ... UK

☐ Mark (X) this box if you attach a continuation sheet.

PART B PARTITION COEFFICIENTS

5.02 a. Specify the half-life of the listed substance in the following media.

<u>Media</u>	<u>Half-life (specify units)</u>
Groundwater	<u>COLIDIFIES ON CONTACT;FORMS NON-TOXIC POLYUREA</u>
Atmosphere	<u>3 HOURS</u>
Surface water	<u>SOLIDIFIES ON CONTACT;FORMS NON-TOXIC POLYUREA</u>
Soil	<u>SOLIDIFIES ON CONTACT;FORMS NON-TOXIC POLYUREA</u>

*SEE ATTACHED INFORMATION

b. Identify the listed substance's known transformation products that have a half-life greater than 24 hours.

<u>CAS No.</u>	<u>Name</u>	<u>Half-life (specify units)</u>	<u>Media</u>
<u>UK</u>	<u></u>	<u></u>	<u>in</u>
<u></u>	<u></u>	<u></u>	<u>in</u>
<u></u>	<u></u>	<u></u>	<u>in</u>
<u></u>	<u></u>	<u></u>	<u>in</u>

5.03 Specify the octanol-water partition coefficient, K_{ow} ... UK at 25°C

Method of calculation or determination

5.04 Specify the soil-water partition coefficient, K_d UK at 25°C

Soil type

5.05 Specify the organic carbon-water partition coefficient, K_{oc} UK at 25°C

5.06 Specify the Henry's Law Constant, H UK atm-m³/mole

☐ Mark (X) this box if you attach a continuation sheet.

5.07 List the bioconcentration factor (BCF) of the listed substance, the species for which it was determined, and the type of test used in deriving the BCF.

<u>Bioconcentration Factor</u>	<u>Species</u>	<u>Test</u> ¹
UK		

¹Use the following codes to designate the type of test:

F = Flowthrough

S = Static

☐ Mark (X) this box if you attach a continuation sheet.

6.04 For each market listed below, state the quantity sold and the total sales value of
CBI the listed substance sold or transferred in bulk during the reporting year.

☐

<u>Market</u>	<u>Quantity Sold or Transferred (kg/yr)</u>	<u>Total Sales Value (\$/yr)</u>
Retail sales	_____	_____
Distribution -- Wholesalers	_____	_____
Distribution -- Retailers	_____	_____
Intra-company transfer	_____	_____
Repackagers	_____	_____
Mixture producers	_____	_____
Article producers	_____	_____
Other chemical manufacturers or processors	_____	_____
Exporters	_____	_____
Other (specify)	_____	_____
_____	_____	_____

6.05 Substitutes -- List all known commercially feasible substitutes that you know exist
for the listed substance and state the cost of each substitute. A commercially
CBI feasible substitute is one which is economically and technologically feasible to use
in your current operation, and which results in a final product with comparable
performance in its end uses.

☐

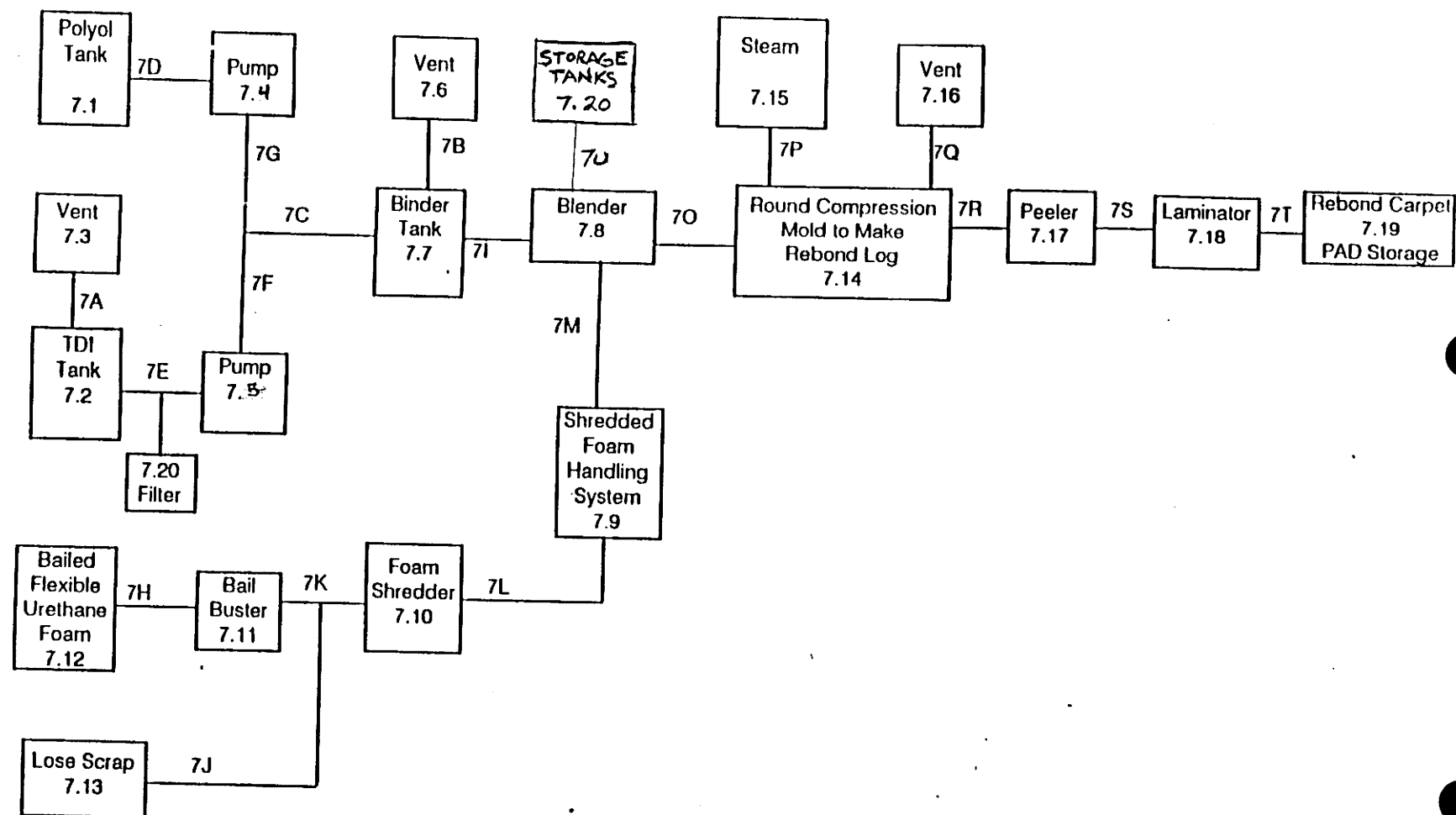
<u>Substitute</u>	<u>Cost (\$/kg)</u>
UK	_____
_____	_____
_____	_____

☐ Mark (X) this box if you attach a continuation sheet.

7.01 MANUFACTURER

Process Type: Rebond Carpet PAD Manufacturing Process

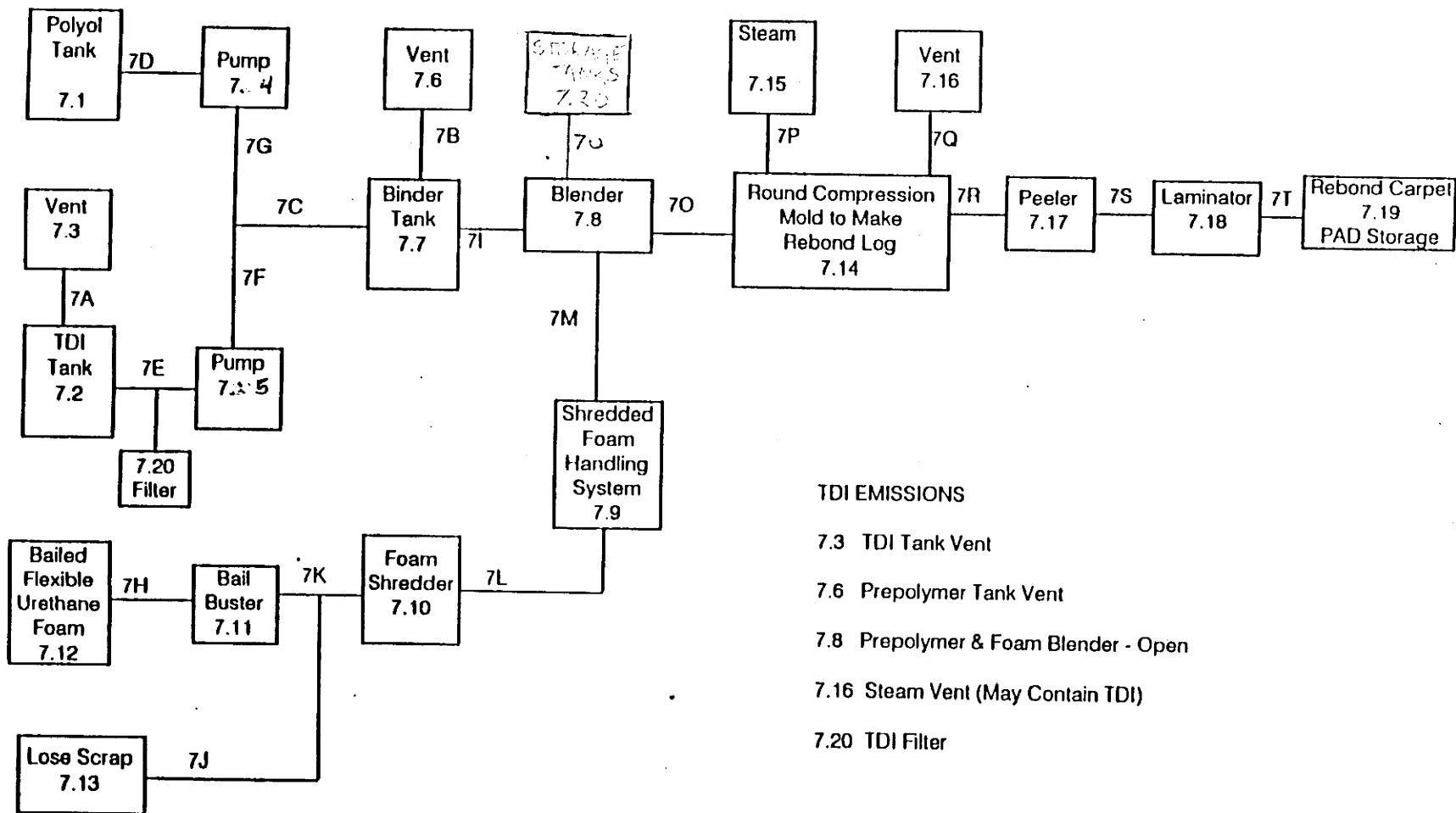
Intermediates: Prepolymer Containing TDI Used to Glue Scrap Foam Into Rebond Log



7.03 EMISSIONS

Process Type: Rebond Foam Carpet PAD Manufacturing Process

Intermediates: Prepolymer Containing TDI Used to Glue Scrap Foam Into Rebond Log



7.04 Describe the typical equipment types for each unit operation identified in your process block flow diagram(s). If a process block flow diagram is provided for more than one process type, photocopy this question and complete it separately for each process type.

CBI

☐ Process type REBOND CARPET MANUFACTURING PROCESS

<u>Unit Operation ID Number</u>	<u>Typical Equipment Type</u>	<u>Operating Temperature Range (°C)</u>	<u>Operating Pressure Range (mm Hg)</u>	<u>Vessel Composition</u>
<u>7.1</u>	<u>STORAGE TANK</u>	<u>32-38°C</u>	<u>760-8500</u>	<u>STEEL</u>
<u>7.2</u>	<u>TDI STORAGE TANK</u>	<u>32-38°C</u>	<u>760-8500</u>	<u>STEEL</u>
<u>7.3</u>	<u>TANK VENT</u>	<u>AMBIENT</u>	<u>760-7000</u>	<u>STEEL</u>
<u>7.4</u>	<u>GEAR PUMP</u>	<u>AMBIENT</u>	<u>760-2070</u>	<u>STEEL</u>
<u>7.5</u>	<u>GEAR PUMP</u>	<u>AMBIENT</u>	<u>760-5700</u>	<u>STEEL</u>
<u>7.6</u>	<u>TANK VENT</u>	<u>AMBIENT</u>	<u>760-7000</u>	<u>STEEL</u>
<u>7.7</u>	<u>BINDER TANK</u>	<u>32-38°C</u>	<u>760-1000</u>	<u>STEEL</u>
<u>7.8</u>	<u>BLENDING TANK</u>	<u>32-38°C</u>	<u>760-1000</u>	<u>STEEL</u>
<u>7.9</u>	<u>STORAGE HOPPERS FOR FOAM PARTICLES</u>	<u>AMBIENT</u>	<u>ATMOSPHERIC</u>	<u>STEEL</u>
<u>7.10</u>	<u>FOAM SHREDDING MACHINES</u>	<u>AMBIENT</u>	<u>ATMOSPHERIC</u>	<u>STEEL</u>

☐ Mark (X) this box if you attach a continuation sheet.

7.04 Describe the typical equipment types for each unit operation identified in your process block flow diagram(s). If a process block flow diagram is provided for more than one process type, photocopy this question and complete it separately for each process type.

CBI

☐ Process type REBOND CARPET MANUFACTURING PROCESS

<u>Unit Operation ID Number</u>	<u>Typical Equipment Type</u>	<u>Operating Temperature Range (°C)</u>	<u>Operating Pressure Range (mm Hg)</u>	<u>Vessel Composition</u>
<u>7.14</u>	<u>MOLDING PROCESS</u>	<u>32-38°C</u>	<u>UK</u>	<u>STEEL</u>
<u>7.15</u>	<u>H2O STEAM INLETS</u>	<u>32-38°C</u>	<u>760-8500</u>	<u>STEEL</u>
<u>7.16</u>	<u>PROCESS VENT</u>	<u>AMBIENT</u>	<u>ATMOSPHERIC</u>	<u>STEEL</u>
<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>

☐ Mark (X) this box if you attach a continuation sheet.

7.05 Describe each process stream identified in your process block flow diagram(s). If a process block flow diagram is provided for more than one process type, photocopy this question and complete it separately for each process type.

CBI

☐ Process type REBOND CARPET MANUFACTURING PROCESS

<u>Process Stream ID Code</u>	<u>Process Stream Description</u>	<u>Physical State¹</u>	<u>Stream Flow (kg/yr)</u>
<u>7E, 7F, 7C, 7I</u>	<u>TOLUENE DIISOCYANATE</u>	<u>OL</u>	<u>254,000</u>
<u>7D, 7G, 7C, 7I</u>	<u>POLYOL RESIN</u>	<u>OL</u>	<u>1,013,000</u>
<u>7U</u>	<u>ORGANIC PIGMENTS</u>	<u>OL</u>	<u>27,600</u>
<u>7U</u>	<u>FLAME RETARDANTS</u>	<u>OL</u>	<u>1,100</u>
<u>7H, 7J, 7K, 7L, 7M</u>	<u>POLYURETHANE FOAM PARTICLES</u>	<u>SO</u>	<u>11,881,000</u>
<u>7O</u>	<u>PREPOLYMER, ORGANIC PIGMENTS, FLAME RETARDENTS, POLYURETHANE FOAM</u>	<u>SO</u>	<u>UK</u>
<u>7P</u>	<u>STEAM</u>	<u>GC</u>	<u>UK</u>
<u>7R</u>	<u>REBOND CARPET UNDERLAY</u>	<u>SO</u>	<u>11,860,000</u>

¹Use the following codes to designate the physical state for each process stream:

GC = Gas (condensable at ambient temperature and pressure)
 GU = Gas (uncondensable at ambient temperature and pressure)
 SO = Solid
 SY = Sludge or slurry
 AL = Aqueous liquid
 OL = Organic liquid
 IL = Immiscible liquid (specify phases, e.g., 90% water, 10% toluene)

☐ Mark (X) this box if you attach a continuation sheet.

7.06 Characterize each process stream identified in your process block flow diagram(s). If a process block flow diagram is provided for more than one process type, photocopy this question and complete it separately for each process type. (Refer to the CBI instructions for further explanation and an example.)

☐ Process type REBOND CARPET MANUFACTURING PROCESS

a.	b.	c.	d.	e.
Process Stream ID Code	Known Compounds ¹	Concentrations ^{2,3} (% or ppm)	Other Expected Compounds	Estimated Concentrations (% or ppm)
<u>7A, 7B</u>	<u>TOLUENE DIISOCYANATE</u>	<u>0.03PPM</u>	<u>NA</u>	<u>NA</u>
	<u>AIR</u>	<u>99.9%</u>	<u>NA</u>	<u>NA</u>
<u>7E, 7F, 7C</u>	<u>TOLUENE DIISOCYANATE</u>	<u>100%</u>	<u>NA</u>	<u>NA</u>
<u>7D, 7G, 7C</u>	<u>POLYOL RESIN</u>	<u>100%</u>	<u>NA</u>	<u>NA</u>

7.06 continued below

☐ Mark (X) this box if you attach a continuation sheet.

7.06 Characterize each process stream identified in your process block flow diagram(s). If a process block flow diagram is provided for more than one process type, photocopy this question and complete it separately for each process type. (Refer to the CBI instructions for further explanation and an example.)

☐ Process type

a.	b.	c.	d.	e.
Process Stream ID Code	Known Compounds ¹	Concentrations ^{2,3} (% or ppm)	Other Expected Compounds	Estimated Concentrations (% or ppm)
7I	POLYOL RESIN	80%	NA	NA
	TOLUENE DIISOCYANATE	20%	NA	NA
7H, 7J, 7K 7L, 7M	POLYURETHANE FOAM	100%	NA	NA
7O	PREPOLYMER, ORGANIC			
	PIGMENTS, FLAME	100%	NA	NA
	RETARDANTS,			
	POLYURETHANE FOAM			

7.06 continued below

☐ Mark (X) this box if you attach a continuation sheet.

7.06 Characterize each process stream identified in your process block flow diagram(s).
 If a process block flow diagram is provided for more than one process type, photocopy
 this question and complete it separately for each process type. (Refer to the
 CBI instructions for further explanation and an example.)

☐ Process type _____

a.	b.	c.	d.	e.
Process Stream ID Code	Known Compounds ¹	Concen- trations ^{2,3} (% or ppm)	Other Expected Compounds	Estimated Concentrations (% or ppm)
7P	H2O STEAM	100%	NA	NA
7Q	TOLUENE DIISOCYANATE	0.04PPM(A/V)	CARBON DIOXIDE	0.1 - 0.5%
	AIR	99.9%		
7R	REBOND CARPET	100%	NA	NA

7.06 continued below

☐ Mark (X) this box if you attach a continuation sheet.

7.06 (continued)

¹For each additive package introduced into a process stream, specify the compounds that are present in each additive package, and the concentration of each component. Assign an additive package number to each additive package and list this number in column b. (Refer to the instructions for further explanation and an example. Refer to the glossary for the definition of additive package.)

Additive Package Number	Components of Additive Package	Concentrations (% or ppm)
<u>1</u>	<u>NA</u>	
<u>2</u>		
<u>3</u>		
<u>4</u>		
<u>5</u>		

²Use the following codes to designate how the concentration was determined:

A = Analytical result
E = Engineering judgement/calculation

³Use the following codes to designate how the concentration was measured:

V = Volume
W = Weight

☐ Mark (X) this box if you attach a continuation sheet.

SECTION 8 RESIDUAL TREATMENT GENERATION, CHARACTERIZATION, TRANSPORTATION, AND
MANAGEMENT

General Instructions:

For questions 8.04-8.06, provide a separate response for each residual treatment block flow diagram provided in question 8.01, 8.02 or 8.03. Identify the process type from which the information is extracted.

For questions 8.05-8.33, the Stream Identification Codes are those process streams listed in either the Section 7 or Section 8 block flow diagrams which contain residuals for each applicable waste management method.

For questions 8.07-8.33, if residuals are combined before they are handled, list those Stream Identification Codes on the same line.

Questions 8.09-8.33 refer to the waste management activities involving the residuals identified in either the Section 7 or Section 8 block flow diagrams. Not all Stream Identification Codes used in the sample answers (e.g., for the incinerator questions) have corresponding process streams identified in the block flow diagram(s). These Stream Identification codes are for illustrative purposes only.

For questions 8.11-8.33, if you have provided the information requested on one of the EPA Office of Solid Waste surveys listed below within the three years prior to your reporting year, you may submit a copy or reasonable facsimile in lieu of answering those questions which the survey addresses. The applicable surveys are: (1) Hazardous Waste Treatment, Storage, Disposal, and Recycling Survey; (2) Hazardous Waste Generator Survey; or (3) Subtitle D Industrial Facility Mail Survey.

☐ Mark (X) this box if you attach a continuation sheet.

8.05 Characterize each process stream identified in your residual treatment block flow diagram(s). If a residual treatment block flow diagram is provided for more than one process type, photocopy this question and complete it separately for each process type. (Refer to the instructions for further explanation and an example.)

[illegible]

☐ Mark (X) this box if you attach a continuation sheet.

8.05 (continued)

¹Use the following codes to designate the type of hazardous waste:

I = Ignitable
C = Corrosive
R = Reactive
E = EP toxic
T = Toxic
H = Acutely hazardous

²Use the following codes to designate the physical state of the residual:

GC = Gas (condensable at ambient temperature and pressure)
→ GU = Gas (uncondensable at ambient temperature and pressure)
SO = Solid
SY = Sludge or slurry
AL = Aqueous liquid
OL = Organic liquid
IL = Immiscible liquid (specify phases, e.g., 90% water, 10% toluene)

8.05 continued below

☐ Mark (X) this box if you attach a continuation sheet.

8.05 (continued)

³For each additive package introduced into a process stream, specify the compounds that are present in each additive package, and the concentration of each component. Assign an additive package number to each additive package and list this number in column d. (Refer to the instructions for further explanation and an example. Refer to the glossary for the definition of additive package.)

<u>Additive Package Number</u>	<u>Components of Additive Package</u>	<u>Concentrations (% or ppm)</u>
<u>1</u>	<u>NA</u>	
<u>2</u>		
<u>3</u>		
<u>4</u>		
<u>5</u>		

⁴Use the following codes to designate how the concentration was determined:

A = Analytical result

E = Engineering judgement/calculation

8.05 continued below

☐ Mark (X) this box if you attach a continuation sheet.

8.05 (continued)

⁵Use the following codes to designate how the concentration was measured:

V = Volume

W = Weight

⁶Specify the analytical test methods used and their detection limits in the table below. Assign a code to each test method used and list those codes in column e.

<u>Code</u>	<u>Method</u>	<u>Detection Limit</u> <u>(± ug/l)</u>
<u>1</u>	NA	
<u>2</u>		
<u>3</u>		
<u>4</u>		
<u>5</u>		
<u>6</u>		

☐ Mark (X) this box if you attach a continuation sheet.

CBI

[illegible]

²Use the codes provided in Exhibit 8-2 to designate the management methods

58

8.22 Describe the combustion chamber design parameters for each of the three largest (by capacity) incinerators that are used on-site to burn the residuals identified in your process block or residual treatment block flow diagram(s).

☐ CBI

Incinerator	Combustion Chamber Temperature (°C)		Location of Temperature Monitor		Residence Time In Combustion Chamber (seconds)	
	Primary	Secondary	Primary	Secondary	Primary	Secondary
1	NA					
2						
3						

Indicate if Office of Solid Waste survey has been submitted in lieu of response by circling the appropriate response.

Yes 1
No 2

8.23 Complete the following table for the three largest (by capacity) incinerators that are used on-site to burn the residuals identified in your process block or residual treatment block flow diagram(s).

☐ CBI

Incinerator	Air Pollution Control Device ¹	Types of Emissions Data Available
1	NA	NA
2		
3		

Indicate if Office of Solid Waste survey has been submitted in lieu of response by circling the appropriate response.

Yes 1
No 2

¹Use the following codes to designate the air pollution control device:

S = Scrubber (include type of scrubber in parenthesis)
E = Electrostatic precipitator
O = Other (specify) _____

☐ Mark (X) this box if you attach a continuation sheet.

SECTION 9 WORKER EXPOSURE

General Instructions:

Questions 9.03-9.25 apply only to those processes and workers involved in manufacturing or processing the listed substance. Do not include workers involved in residual waste treatment unless they are involved in this treatment process on a regular basis (i.e., exclude maintenance workers, construction workers, etc.).

☐ Mark (X) this box if you attach a continuation sheet.

PART A EMPLOYMENT AND POTENTIAL EXPOSURE PROFILE

9.01 Mark (X) the appropriate column to indicate whether your company maintains records on the following data elements for hourly and salaried workers. Specify for each data element the year in which you began maintaining records and the number of years the records for that data element are maintained. (Refer to the instructions for further explanation and an example.)

CBI

☐

Data Element	Data are Maintained for:		Year in Which Data Collection Began	Number of Years Records Are Maintained
	Hourly Workers	Salaried Workers		
Date of hire	X	X	1986	25
Age at hire	X	X	1986	25
Work history of individual before employment at your facility	X	X	1986	25
Sex	X	X	1986	25
Race	X	X	1986	25
Job titles	X	X	1986	25
Start date for each job title	X	X	1986	25
End date for each job title	X	X	1986	25
Work area industrial hygiene monitoring data	X	X	1986	25
Personal employee monitoring data	NA	NA	NA	NA
Employee medical history	X	X	1986	25
Employee smoking history	NA	NA	1986	25
Accident history	X	X	1986	25
Retirement date	X	X	1986	25
Termination date	X	X	1986	25
Vital status of retirees	NA	NA	NA	NA
Cause of death data	NA	NA	NA	NA

☐ Mark (X) this box if you attach a continuation sheet.

9.02 In accordance with the instructions, complete the following table for each activity in which you engage.

CBI

☐

a.	b.	c.	d.	e.
<u>Activity</u>	<u>Process Category</u>	<u>Yearly Quantity (kg)</u>	<u>Total Workers</u>	<u>Total Worker-Hours</u>
Manufacture of the listed substance	Enclosed	NA	NA	NA
	Controlled Release	NA	NA	NA
	Open	NA	NA	NA
On-site use as reactant	Enclosed	NA	NA	NA
	Controlled Release	254,000	27	46,080
	Open	NA	NA	NA
On-site use as nonreactant	Enclosed	NA	NA	NA
	Controlled Release	NA	NA	NA
	Open	NA	NA	NA
On-site preparation of products	Enclosed	NA	NA	NA
	Controlled Release	NA	NA	NA
	Open	NA	NA	NA

☐

Mark (X) this box if you attach a continuation sheet.

9.03 Provide a descriptive job title for each labor category at your facility that encompasses workers who may potentially come in contact with or be exposed to the listed substance.

CBI

☐

Labor Category

Descriptive Job Title

A

FOREMAN/ASSISTANT FOREMAN

B

MOLDING MACHINE OPERATORS

C

FORKLIFT OPERATORS

D

GRINDER PERSONNEL

E

MAINTENANCE PERSONNEL

F

G

H

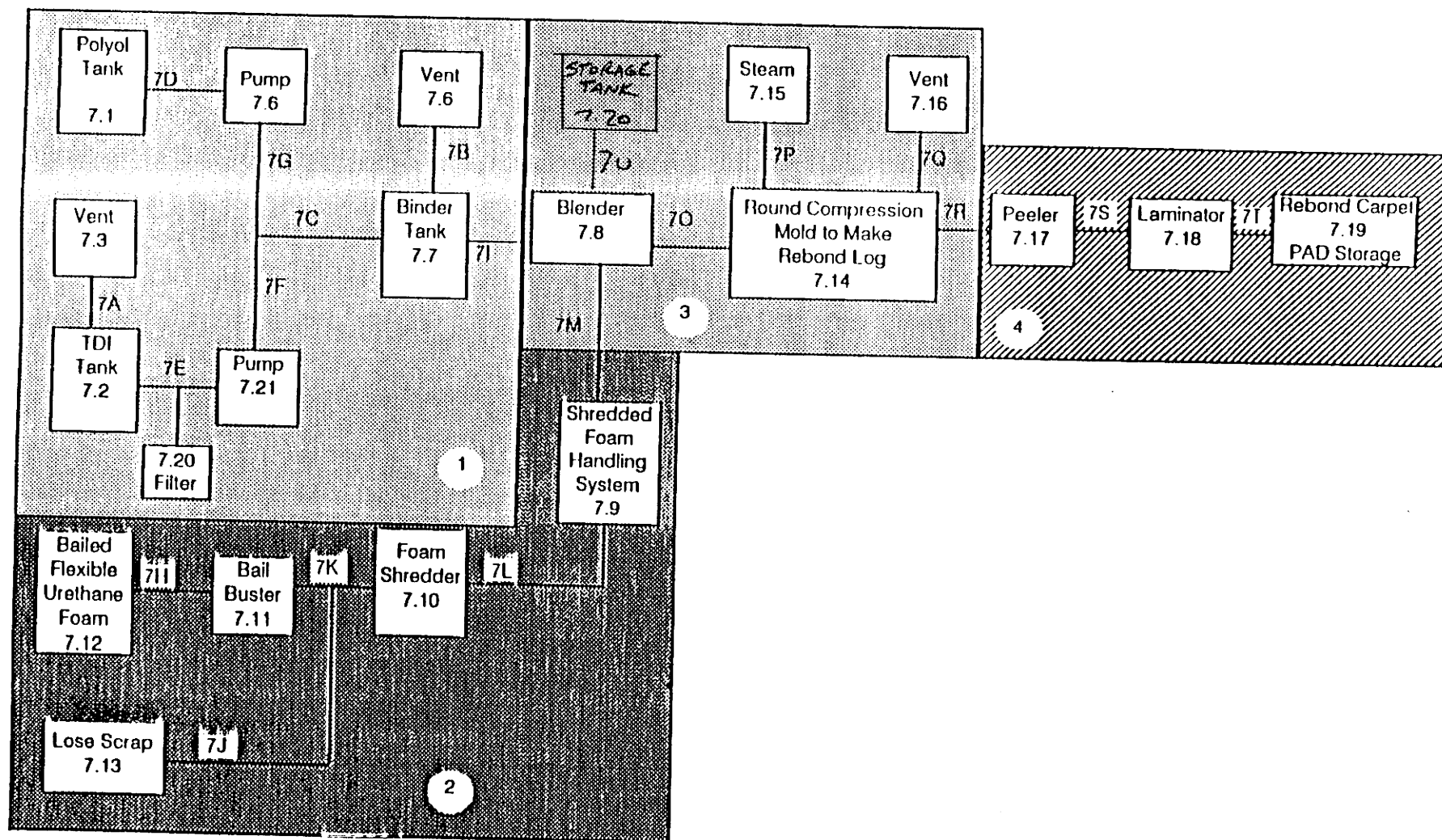
I

J

☐ Mark (X) this box if you attach a continuation sheet.

Process Type: Rebond Carpet PAD Manufacturing Process

Intermediates: Prepolymer Containing TDI Used to Glue Scrap Foam into Rebond Log



9.05 Describe the various work area(s) shown in question 9.04 that encompass workers who may potentially come in contact with or be exposed to the listed substance. Add any additional areas not shown in the process block flow diagram in question 7.01 or 7.02. Photocopy this question and complete it separately for each process type.

CBI

☐ Process type REBOND CARPET MANUFACTURING PROCESS

Work Area ID

Description of Work Areas and Worker Activities

1	<u>STORAGE TANK AREA, PUMPING SYSTEMS, BINDER TANK</u>
2	<u>FOAM SHREDDERS AND STORAGE HOPPERS</u>
3	<u>BLENDER SYSTEM, COMPRESSION MOLDING PROCESS, STEAM SYSTEMS</u>
4	<u>PEELERS, LAMINATORS, STORAGE AREAS</u>
5	<u></u>
6	<u></u>
7	<u></u>
8	<u></u>
9	<u></u>
10	<u></u>

☐ Mark (X) this box if you attach a continuation sheet.

9.06 Complete the following table for each work area identified in question 9.05, and for each labor category at your facility that encompasses workers who may potentially come in contact with or be exposed to the listed substance. Photocopy this question and complete it separately for each process type and work area.

☐ CBI Process type REBOND CARPET MANUFACTURING PROCESS

Work area 1

Labor Category	Number of Workers Exposed	Mode of Exposure (e.g., direct skin contact)	Physical State of Listed Substance ¹	Average Length of Exposure Per Day ²	Number of Days per Year Exposed
E	4	INHALATION	GU	C	50
E	4	DIRECT SKIN CONTACT	OL	NA	NA
A	3	INHALATION	GU	E	260
A	3	DIRECT SKIN CONTACT	OL	NA	NA
A	2	INHALATION	GU	C	50
A	2	DIRECT SKIN CONTACT	OL	NA	NA

¹Use the following codes to designate the physical state of the listed substance at the point of exposure:

GC = Gas (condensable at ambient temperature and pressure)
 GU = Gas (uncondensable at ambient temperature and pressure; includes fumes, vapors, etc.)
 SO = Solid

SY = Sludge or slurry
 AL = Aqueous liquid
 OL = Organic liquid
 IL = Immiscible liquid (specify phases, e.g., 90% water, 10% toluene)

²Use the following codes to designate average length of exposure per day:

A = 15 minutes or less
 B = Greater than 15 minutes, but not exceeding 1 hour
 C = Greater than one hour, but not exceeding 2 hours

D = Greater than 2 hours, but not exceeding 4 hours
 E = Greater than 4 hours, but not exceeding 8 hours
 F = Greater than 8 hours

☐ Mark (X) this box if you attach a continuation sheet.

9.06 Complete the following table for each work area identified in question 9.05, and for each labor category at your facility that encompasses workers who may potentially come in contact with or be exposed to the listed substance. Photocopy this question and complete it separately for each process type and work area.

☐ Process type REBOND CARPET MANUFACTURING PROCESS

Work area 2

Labor Category	Number of Workers Exposed	Mode of Exposure (e.g., direct skin contact)	Physical State of Listed Substance ¹	Average Length of Exposure Per Day ²	Number of Days per Year Exposed
D	9	INHALATION	GU	E	260
C	6	INHALATION	GU	E	260

¹Use the following codes to designate the physical state of the listed substance at the point of exposure:

GC = Gas (condensable at ambient temperature and pressure)
 GU = Gas (uncondensable at ambient temperature and pressure; includes fumes, vapors, etc.)
 SO = Solid

SY = Sludge or slurry
 AL = Aqueous liquid
 OL = Organic liquid
 IL = Immiscible liquid (specify phases, e.g., 90% water, 10% toluene)

²Use the following codes to designate average length of exposure per day:

A = 15 minutes or less
 B = Greater than 15 minutes, but not exceeding 1 hour
 C = Greater than one hour, but not exceeding 2 hours

D = Greater than 2 hours, but not exceeding 4 hours
 E = Greater than 4 hours, but not exceeding 8 hours
 F = Greater than 8 hours

☐ Mark (X) this box if you attach a continuation sheet.

9.06 Complete the following table for each work area identified in question 9.05, and for each labor category at your facility that encompasses workers who may potentially come in contact with or be exposed to the listed substance. Photocopy this question and complete it separately for each process type and work area.

☐ Process type REBOND CARPET MANUFACTURING PROCESS

Work area 3

Labor Category	Number of Workers Exposed	Mode of Exposure (e.g., direct skin contact)	Physical State of Listed Substance ¹	Average Length of Exposure Per Day ²	Number of Days per Year Exposed
B	3	INHALATION	GU	E	260

¹Use the following codes to designate the physical state of the listed substance at the point of exposure:

GC = Gas (condensable at ambient temperature and pressure)	SY = Sludge or slurry
GU = Gas (uncondensable at ambient temperature and pressure; includes fumes, vapors, etc.)	AL = Aqueous liquid
SO = Solid	OL = Organic liquid
	IL = Immiscible liquid (specify phases, e.g., 90% water, 10% toluene)

²Use the following codes to designate average length of exposure per day:

A = 15 minutes or less	D = Greater than 2 hours, but not exceeding 4 hours
B = Greater than 15 minutes, but not exceeding 1 hour	E = Greater than 4 hours, but not exceeding 8 hours
C = Greater than one hour, but not exceeding 2 hours	F = Greater than 8 hours

☐ Mark (X) this box if you attach a continuation sheet.

9.07 For each labor category represented in question 9.06, indicate the 8-hour Time Weighted Average (TWA) exposure levels and the 15-minute peak exposure levels. Photocopy this question and complete it separately for each process type and work area.

CBI

☐ Process type REBOND CARPET MANUFACTURING PROCESS

Work area 1-3

Labor Category	8-hour TWA Exposure Level (ppm, mg/m ³ , other-specify)	15-Minute Peak Exposure Level (ppm, mg/m ³ , other-specify)
A-E	<1.0 PPB	<1.0 PPB

☐ Mark (X) this box if you attach a continuation sheet.

PART B WORK PLACE MONITORING PROGRAM

9.08 If you monitor worker exposure to the listed substance, complete the following table.

CBI

☐

<u>Sample/Test</u>	<u>Work Area ID</u>	<u>Testing Frequency (per year)</u>	<u>Number of Samples (per test)</u>	<u>Who Samples¹</u>	<u>Analyzed In-House (Y/N)</u>	<u>Number of Years Records Maintained</u>
Personal breathing zone	NA	NA	NA	NA	NA	NA
General work area (air)	1-4	OCCASIONAL	CONTINUOUS	D	Y	5 (MINIMUM)
Wipe samples	NA	NA	NA	NA	NA	NA
Adhesive patches	NA	NA	NA	NA	NA	NA
Blood samples	NA	NA	NA	NA	NA	NA
Urine samples	NA	NA	NA	NA	NA	NA
Respiratory samples	NA	NA	NA	NA	NA	NA
Allergy tests	NA	NA	NA	NA	NA	NA
Other (specify)						
Other (specify)						
Other (specify)						

¹Use the following codes to designate who takes the monitoring samples:

- A = Plant industrial hygienist
- B = Insurance carrier
- C = OSHA consultant
- D = Other (specify) PLANT PERSONNEL

☐ Mark (X) this box if you attach a continuation sheet.

9.09 For each sample type identified in question 9.08, describe the type of sampling and analytical methodology used for each type of sample.

☐ Sample Type Sampling and Analytical Methodology

GENERAL WORK AREA STATIONARY AND PORTABLE MONITORS

9.10 If you conduct personal and/or ambient air monitoring for the listed substance, specify the following information for each equipment type used.

CBI

<input type="checkbox"/> <u>Equipment Type</u> ¹	<u>Detection Limit</u> ²	<u>Manufacturer</u>	<u>Averaging Time (hr)</u>	<u>Model Number</u>
E	<.001A	MDA SCIENTIFIC	8	7005
H	<.001A	GMD SYSTEMS	0.15	900

¹Use the following codes to designate personal air monitoring equipment types:

- A = Passive dosimeter
- B = Detector tube
- C = Charcoal filtration tube with pump
- D = Other (specify) _____

Use the following codes to designate ambient air monitoring equipment types:

- E = Stationary monitors located within work area
- F = Stationary monitors located within facility
- G = Stationary monitors located at plant boundary
- H = Mobile monitoring equipment (specify) PORTABLE AUTO-STEP MONITOR
- I = Other (specify) _____

²Use the following codes to designate detection limit units:

- A = ppm
- B = Fibers/cubic centimeter (f/cc)
- C = Micrograms/cubic meter (μ/m^3)

☐ Mark (X) this box if you attach a continuation sheet.

9.11 If you conduct routine medical tests for monitoring the health effects of exposure to the listed substance, specify the type and frequency of the tests.

CBI

☐

Test Description

Frequency
(weekly, monthly, yearly, etc.)

PULIMONARY FUNCTION STUDIES

VOLUNTARY BASIS

☐ Mark (X) this box if you attach a continuation sheet.

9.12 Describe the engineering controls that you use to reduce or eliminate worker exposure to the listed substance. Photocopy this question and complete it separately for each process type and work area.

☐ Process type REBOND CARPET MANUFACTURING PROCESS

Work area 3

<u>Engineering Controls</u>	<u>Used (Y/N)</u>	<u>Year Installed</u>	<u>Upgraded (Y/N)</u>	<u>Year Upgraded</u>
Ventilation:				
Local exhaust	<u>Y</u>	<u>1986</u>	<u>Y</u>	<u>1988</u>
General dilution	<u>Y</u>	<u>1986</u>	<u>Y</u>	<u>1987</u>
Other (specify) <u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
Vessel emission controls	<u>Y</u>	<u>1987</u>	<u>NA</u>	<u>NA</u>
Mechanical loading or packaging equipment	<u>N</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>
Other (specify) <u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>

98

9.13 Describe all equipment or process modifications you have made within the 3 years prior to the reporting year that have resulted in a reduction of worker exposure to the listed substance. For each equipment or process modification described, state the percentage reduction in exposure that resulted. Photocopy this question and complete it separately for each process type and work area.

CBI

☐ Process type REBOND CARPET MANUFACTURING PROCESS

Work area 3

<u>Equipment or Process Modification</u>	<u>Reduction in Worker Exposure Per Year (%)</u>
<u>COMPUTERIZED MOLD PROCESS FROM MANUAL BASIS</u>	<u>NA</u>
<u>MOLD EXHAUST PIPING INSTALLED TO OUTSIDE EXHAUST FANS</u>	<u>NA</u>
<u>INSTALLED 3 ADDITIONAL OUTSIDE EXHAUST FANS</u>	<u>NA</u>
<u>PURCHASED MOLD CLEANER TO CLEAN INSIDE OF MOLDS AUTOMATICALLY.</u>	<u>NA</u>

☐ Mark (X) this box if you attach a continuation sheet.

PART D PERSONAL PROTECTIVE AND SAFETY EQUIPMENT

9.14 Describe the personal protective and safety equipment that your workers wear or use in each work area in order to reduce or eliminate their exposure to the listed substance. Photocopy this question and complete it separately for each process type and work area.

CBI

[] Process type REBOND CARPET MANUFACTURING PROCESS

Work area 3

<u>Equipment Types</u>	<u>Wear or Use (Y/N)</u>
Respirators	<u>Y (1)</u>
Safety goggles/glasses	<u>Y (1)</u>
Face shields	<u>N</u>
Coveralls	<u>N</u>
Bib aprons	<u>N</u>
Chemical-resistant gloves	<u>Y (1)</u>
Other (specify)	
<u>SELF CONTAINED</u>	<u></u>
<u>BREATHING APPARATUS</u>	<u>Y (1)</u>

(1) NOT USED DURING NORMAL PRODUCTION CYCLE. USED BASICALLY WHEN IN CONFINED AREA WHEN DEALING WITH THE HAZARDOUS CHEMICAL TO REPAIR OR REPLACE MALFUNCTIONING EQUIPMENT.

[] Mark (X) this box if you attach a continuation sheet.

9.15 If workers use respirators when working with the listed substance, specify for each process type, the work areas where the respirators are used, the type of respirators used, the average usage, whether or not the respirators were fit tested, and the type and frequency of the fit tests. Photocopy this question and complete it separately for each process type.

CBI

☐ Process type REBOND CARPET MANUFACTURING PROCESS

Work Area	Respirator Type	Average Usage ¹	Fit Tested (Y/N)	Type of Fit Test ²	Frequency of Fit Tests (per year)
<u>3</u>	<u>COMFO II FACE PIECE</u>	<u>E</u>	<u>N</u>	<u>NA</u>	<u>NA</u>
_____	_____	_____	_____	_____	_____
_____	_____	_____	_____	_____	_____
_____	_____	_____	_____	_____	_____

¹Use the following codes to designate average usage:

A = Daily
 B = Weekly
 C = Monthly
 D = Once a year
 E = Other (specify) WHEN NECESSARY

²Use the following codes to designate the type of fit test:

QL = Qualitative
 QT = Quantitative

☐ Mark (X) this box if you attach a continuation sheet.

PART E WORK PRACTICES

- 9.19 Describe all of the work practices and administrative controls used to reduce or eliminate worker exposure to the listed substance (e.g., restrict entrance only to authorized workers, mark areas with warning signs, insure worker detection and monitoring practices, provide worker training programs, etc.). Photocopy this question and complete it separately for each process type and work area.

CBI

☐

Process type REBOND CARPET MANUFACTURING PROCESS

Work area 1-3

EXPOSURE MONITORING, RESPIRATOR PROTECTION, TRAINING PROGRAM, SELF
CONTAINED BREATHING APPARATUS, LIMITED ACCESS DIKING, NEUTRALIZER
AND SPILL ABSORBANT, TRAINING FILMS

- 9.20 Indicate (X) how often you perform each housekeeping task used to clean up routine leaks or spills of the listed substance. Photocopy this question and complete it separately for each process type and work area.

Process type REBOND CARPET MANUFACTURING PROCESS

Work area 1-3

<u>Housekeeping Tasks</u>	<u>Less Than Once Per Day</u>	<u>1-2 Times Per Day</u>	<u>3-4 Times Per Day</u>	<u>More Than 4 Times Per Day</u>
Sweeping	<u> </u>	<u> </u>	<u> X </u>	<u> </u>
Vacuuming	<u> </u>	<u> </u>	<u> </u>	<u> </u>
Water flushing of floors	<u> </u>	<u> </u>	<u> </u>	<u> </u>
Other (specify)				
<u>EMERGENCY-IMMEDIATE CLEAN UP, DIKING, ABSORBANT</u>				

☐ Mark (X) this box if you attach a continuation sheet.

9.21 Do you have a written medical action plan for responding to routine or emergency exposure to the listed substance?

Routine exposure

Yes (1)
No 2

Emergency exposure

Yes (1)
No 2

If yes, where are copies of the plan maintained?

Routine exposure: PLANT MANAGERS OFFICE

Emergency exposure: SAME AS ABOVE

9.22 Do you have a written leak and spill cleanup plan that addresses the listed substance? Circle the appropriate response.

Yes (1)
No 2

If yes, where are copies of the plan maintained? PLANT MANAGERS OFFICE

Has this plan been coordinated with state or local government response organizations?
Circle the appropriate response.

Yes (1)
No 2

9.23 Who is responsible for monitoring worker safety at your facility? Circle the appropriate response.

Plant safety specialist 1
Insurance carrier 2
OSHA consultant 3
Other (specify) R&D DEPARTMENT/PLANT SUPERVISION (4)

[] Mark (X) this box if you attach a continuation sheet.

SECTION 10 ENVIRONMENTAL RELEASE

General Instructions:

Complete Part E (questions 10.23-10.35) for each non-routine release involving the listed substance that occurred during the reporting year. Report on all releases that are equal to or greater than the listed substance's reportable quantity value, RQ, unless the release is federally permitted as defined in 42 U.S.C. 9601, or is specifically excluded under the definition of release as defined in 40 CFR 302.3(22). Reportable quantities are codified in 40 CFR Part 302. If the listed substance is not a hazardous substance under the Comprehensive Environmental Response, Compensation, and Liability Act of 1980 (CERCLA) and, thus, does not have an RQ, then report releases that exceed 2,270 kg. If such a substance however, is designated as a CERCLA hazardous substance, then report those releases that are equal to or greater than the RQ. The facility may have answered these questions or similar questions under the Agency's Accidental Release Information Program and may already have this information readily available. Assign a number to each release and use this number throughout this part to identify the release. Releases over more than a 24-hour period are not single releases, i.e., the release of a chemical substance equal to or greater than an RQ must be reported as a separate release for each 24-hour period the release exceeds the RQ.

For questions 10.25-10.35, answer the questions for each release identified in question 10.23. Photocopy these questions and complete them separately for each release.

PART A GENERAL INFORMATION

10.01 Where is your facility located? Circle all appropriate responses.

CBI

- ☐ Industrial area 1
- Urban area 2
- Residential area 3
- Agricultural area 4
- Rural area 5
- Adjacent to a park or a recreational area 6
- Within 1 mile of a navigable waterway 7
- Within 1 mile of a school, university, hospital, or nursing home facility 8
- Within 1 mile of a non-navigable waterway 9
- Other (specify) NUCLEAR POWER PLANT 10

☐ Mark (X) this box if you attach a continuation sheet.

10.02 Specify the exact location of your facility (from central point where process unit is located) in terms of latitude and longitude or Universal Transverse Mercader (UTM) coordinates.

Latitude 41 ° 07 ' 00 "

Longitude 76 ° 08 ' 30 "

UTM coordinates Zone UK , Northing UK , Easting UK

10.03 If you monitor meteorological conditions in the vicinity of your facility, provide the following information.

Average annual precipitation inches/year

Predominant wind direction

10.04 Indicate the depth to groundwater below your facility.

Depth to groundwater meters

10.05 For each on-site activity listed, indicate (Y/N/NA) all routine releases of the listed substance to the environment. (Refer to the instructions for a definition of CBI Y, N, and NA.)

<input type="checkbox"/> On-Site Activity	Environmental Release		
	Air	Water	Land
Manufacturing	NA	NA	NA
Importing	NA	NA	NA
Processing	Y	N	N
Otherwise used	NA	NA	NA
Product or residual storage	Y	N	N
Disposal	NA	NA	NA
Transport	NA	NA	NA

☐ Mark (X) this box if you attach a continuation sheet.

10.06 Provide the following information for the listed substance and specify the level of precision for each item. (Refer to the instructions for further explanation and an example.)

CBI

☐

Quantity discharged to the air 40 kg/yr \pm 20 %

Quantity discharged in wastewaters NA kg/yr \pm %

Quantity managed as other waste in on-site
treatment, storage, or disposal units NA kg/yr \pm %

Quantity managed as other waste in off-site
treatment, storage, or disposal units NA kg/yr \pm %

☐ Mark (X) this box if you attach a continuation sheet.

10.08 Describe the control technologies used to minimize release of the listed substance for each process stream containing the listed substance as identified in your process block or residual treatment block flow diagram(s). Photocopy this question and complete it separately for each process type.

CBI

☐ Process type REBOND CARPET MANUFACTURING PROCESS

<u>Stream ID Code</u>	<u>Control Technology</u>	<u>Percent Efficiency</u>
<u>NA</u>		

☐ Mark (X) this box if you attach a continuation sheet.

PART B RELEASE TO AIR

- 10.09 Point Source Emissions -- Identify each emission point source containing the listed substance in terms of a Stream ID Code as identified in your process block or residual treatment block flow diagram(s), and provide a description of each point source. Do not include raw material and product storage vents, or fugitive emission sources (e.g., equipment leaks). Photocopy this question and complete it separately for each process type.

CBI

☐

Process type REBOND CARPET MANUFACTURING PROCESS

Point Source
ID Code

Description of Emission Point Source

7A

STORAGE TANK VENT

7B

BINDER TANK VENT

7Q

PROCESS VENT STACKS

☐ Mark (X) this box if you attach a continuation sheet.

☐ Mark (X) this box if you attach a continuation sheet.

10.10 Emission Characteristics -- Characterize the emissions for each Point Source ID Code identified in question 10.09 by completing the following table.

CBI

<input type="checkbox"/> Point Source ID Code	Physical State ¹	Average Emissions (kg/day)	Frequency ² (days/yr)	Duration ³ (min/day)	Average Emission Factor ⁴	Maximum Emission Rate (kg/min)	Maximum Emission Rate Frequency (events/yr)	Maximum Emission Rate Duration (min/event)
7Q, 7B	V	0.15	260	927	NA	.00015	68,900	3.5
7A	V	0.003	15	120	NA	NA	15	120

¹Use the following codes to designate physical state at the point of release:

G = Gas; V = Vapor; P = Particulate; A = Aerosol; O = Other (specify) _____

²Frequency of emission at any level of emission

³Duration of emission at any level of emission

⁴Average Emission Factor -- Provide estimated (\pm 25 percent) emission factor (kg of emission per kg of production of listed substance)

10.11 Stack Parameters -- Identify the stack parameters for each Point Source ID Code identified in question 10.09 by completing the following table.

CBI

☐

Point Source ID Code	Stack Height(m)	Stack Inner Diameter (at outlet) (m)	Exhaust Temperature (°C)	Emission Exit Velocity (m/sec)	Building Height(m) ¹	Building Width(m) ²	Vent Type ³
7Q	2M	.9M	AMBIENT	12.5	10M	32M	V

¹Height of attached or adjacent building

²Width of attached or adjacent building

³Use the following codes to designate vent type:

H = Horizontal

V = Vertical

☐ Mark (X) this box if you attach a continuation sheet.

10.12 If the listed substance is emitted in particulate form, indicate the particle size distribution for each Point Source ID Code identified in question 10.09.
Photocopy this question and complete it separately for each emission point source.

CBI

☐

Point source ID code NA

Size Range (microns)

Mass Fraction (% \pm % precision)

< 1	<u>NA</u>
≥ 1 to < 10	<u>NA</u>
≥ 10 to < 30	<u>NA</u>
≥ 30 to < 50	<u>NA</u>
≥ 50 to < 100	<u>NA</u>
≥ 100 to < 500	<u>NA</u>
≥ 500	<u>NA</u>

Total = 100%

☐ Mark (X) this box if you attach a continuation sheet.

PART C FUGITIVE EMISSIONS

10.13 Equipment Leaks -- Complete the following table by providing the number of equipment types listed which are exposed to the listed substance and which are in service according to the specified weight percent of the listed substance passing through the component. Do this for each process type identified in your process block or residual treatment block flow diagram(s). Do not include equipment types that are not exposed to the listed substance. If this is a batch or intermittently operated process, give an overall percentage of time per year that the process type is exposed to the listed substance. Photocopy this question and complete it separately for each process type.

CBI

☐ Process type REBOND CARPET MANUFACTURING PROCESS

Percentage of time per year that the listed substance is exposed to this process type 100 %

Equipment Type	Number of Components in Service by Weight Percent of Listed Substance in Process Stream					Greater than 99%
	Less than 5%	5-10%	11-25%	26-75%	76-99%	
Pump seals ¹						
Packed						
Mechanical			2			1
Double mechanical ²						
Compressor seals ¹						
Flanges	12		18			6
Valves						
Gas ³						
Liquid	2		6			3
Pressure relief devices ⁴ (Gas or vapor only)						
Sample connections						
Gas						
Liquid						
Open-ended lines ⁵ (e.g., purge, vent)						
Gas	1					
Liquid						

¹List the number of pump and compressor seals, rather than the number of pumps or compressors

10.13 continued on next page

☐ Mark (X) this box if you attach a continuation sheet.

10.13 (continued)

²If double mechanical seals are operated with the barrier (B) fluid at a pressure greater than the pump stuffing box pressure and/or equipped with a sensor (S) that will detect failure of the seal system, the barrier fluid system, or both, indicate with a "B" and/or an "S", respectively

³Conditions existing in the valve during normal operation

⁴Report all pressure relief devices in service, including those equipped with control devices

⁵Lines closed during normal operation that would be used during maintenance operations

10.14 Pressure Relief Devices with Controls -- Complete the following table for those pressure relief devices identified in 10.13 to indicate which pressure relief devices in service are controlled. If a pressure relief device is not controlled, enter "None" under column c.

☐

a. Number of Pressure Relief Devices	b. Percent Chemical in Vessel ¹	c. Control Device	d. Estimated Control Efficiency ²
1	100%	PRESSURE RELIEF	100%

¹Refer to the table in question 10.13 and record the percent range given under the heading entitled "Number of Components in Service by Weight Percent of Listed Substance" (e.g., <5%, 5-10%, 11-25%, etc.)

²The EPA assigns a control efficiency of 100 percent for equipment leaks controlled with rupture discs under normal operating conditions. The EPA assigns a control efficiency of 98 percent for emissions routed to a flare under normal operating conditions

☐ Mark (X) this box if you attach a continuation sheet.

10.15 Equipment Leak Detection -- If a formal leak detection and repair program is in place, complete the following table regarding those leak detection and repair procedures. Photocopy this question and complete it separately for each process type.

CBI

☐ Process typeREBOND CARPET MANUFACTURING PROCESS

<u>Equipment Type</u>	<u>Leak Detection</u> <u>Concentration</u> (ppm or mg/m ³)	<u>Detection</u> <u>Device</u> ¹	<u>Frequency</u> <u>of Leak</u> <u>Detection</u> (per year)	<u>Repairs</u> <u>Initiated</u> (days after detection)	<u>Repairs</u> <u>Completed</u> (days after initiated)
	Measured at Inches from Source				
Pump seals					
Packed	NA				
Mechanical					
Double mechanical					
Compressor seals					
Flanges					
Valves					
Gas					
Liquid					
Pressure relief devices (gas or vapor only)					
Sample connections					
Gas					
Liquid					
Open-ended lines					
Gas					
Liquid					

¹Use the following codes to designate detection device:

POVA = Portable organic vapor analyzer

FPM = Fixed point monitoring

0 = Other (specify) _____

☐ Mark (X) this box if you attach a continuation sheet.

☐ Mark (X) this box if you attach a continuation sheet.

10.16 Raw Material, Intermediate and Product Storage Emissions - - Complete the following table by providing the information on each liquid raw material, intermediate, and product storage vessel containing the listed substance as identified in your process block or residual treatment block flow diagram(s).

Vessel Type ¹	Floating Roof ² Seals	Composition of Stored Materials ³	Throughput (liters per year)	Vessel Filling Rate	Vessel Filling Duration	Vessel Inner Diameter	Vessel Height	Operating Volume	Vessel Emission Controls ⁴	Design Flow Rate ⁵	Vent Diameter	Control Efficiency	Basis for Estimate ⁶
				(gpm)	(min)	(m)	(m)	(l)			(cm)	(%)	
H	NA	100%	UK	60	90	3.5	2.5	18,900	NA	NA	8.5	NA	NA

¹Use the following codes to designate vessel type:

F = Fixed roof
 CIF = Contact internal floating roof
 NCIF = Noncontact internal floating roof
 EFR = External floating roof
 P = Pressure vessel (indicate pressure rating)
 H = Horizontal
 U = Underground

²Use the following codes to designate floating roof seals:

MS1 = Mechanical shoe, primary
 MS2 = Shoe-mounted secondary
 MS2R = Rim-mounted, secondary
 LM1 = Liquid-mounted resilient filled seal, primary
 LM2 = Rim-mounted shield
 LMW = Weather shield
 VM1 = Vapor mounted resilient filled seal, primary
 VM2 = Rim-mounted secondary
 VMW = Weather shield

³Indicate weight percent of the listed substance. Include the total volatile organic content in parenthesis

⁴Other than floating roofs

⁵Gas/vapor flow rate the emission control device was designed to handle (specify flow rate units)

⁶Use the following codes to designate basis for estimate of control efficiency:

C = Calculations
 S = Sampling

PART E NON-ROUTINE RELEASES

10.23 Indicate the date and time when the release occurred and when the release ceased or was stopped. If there were more than six releases, attach a continuation sheet and list all releases.

<u>Release</u>	<u>Date Started</u>	<u>Time (am/pm)</u>	<u>Date Stopped</u>	<u>Time (am/pm)</u>
<u>1</u>	<u>NA</u>	<u> </u>	<u> </u>	<u> </u>
<u>2</u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u>3</u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u>4</u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u>5</u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u>6</u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>

10.24 Specify the weather conditions at the time of each release.

<u>Release</u>	<u>Wind Speed (km/hr)</u>	<u>Wind Direction</u>	<u>Humidity (%)</u>	<u>Temperature (°C)</u>	<u>Precipitation (Y/N)</u>
<u>1</u>	<u>NA</u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u>2</u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u>3</u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u>4</u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u>5</u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>
<u>6</u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>

☐ Mark (X) this box if you attach a continuation sheet.

Fate of TDI and MDI in Air, Soil, and Water

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ABSTRACT

Toluene diisocyanate (TDI) and methylene diphenylene diisocyanate (MDI) are used in the production of polyurethanes. They can cause respiratory problems at very low concentrations, and workplace and emission levels have been subject to rigorous controls for many years. As a result of these controls, and the very low vapour pressures of the products and their variants, environmental pollution due to emissions or spillages is very low.

III has sponsored a range of studies to determine the fate of TDI and MDI in air, soil and water. Studies of simulated atmospheric conditions indicate that TDI is destroyed predominantly by OH radicals, without the formation of toluene diamine (TDA). TDA or MDA (methylene dianiline), if generated in the atmosphere from any source, are also destroyed by OH radical attack, and no accumulation of these products is expected.

In soil and water TDI and MDI are converted to polyureas, which are chemically inert, and which appear to cause no toxicological effects. The initial rate of reaction of TDI and MDI with water is relatively fast, but in many conditions the resulting polyurea products encapsulate agglomerations of the diisocyanates and rates of reactions decrease rapidly. Under aquatic conditions TDA and MDA are produced in low, transient, concentrations. Studies of the interactions of TDI and MDI with bioaquatic systems are difficult to execute consistently, due to the problem of formulating and controlling suitable conditions of chemical addition. However, the broad overview is that the ecological impact of such interactions is likely to be slight and reversible. III continues its work in these fields.

INTRODUCTION

Polyurethanes are remarkable materials which are used in many aspects of modern life, including insula-

tion, furnishing, construction, surface coatings, sport and medical care. In recent years a range of diisocyanates have been introduced in the manufacture of polyurethanes, but toluene diisocyanate (TDI) and methylene diphenylene diisocyanate (MDI) still dominate the field. World production of each is currently approaching 1 million tons per annum. The International Isocyanate Institute, Inc. (III) is an association of manufacturers of TDI and MDI, and its Member Companies produce a very large proportion of total world tonnage. The main aim of III is the promotion of the safe handling of TDI and MDI, and it has made a major contribution to our knowledge of the environmental effects of TDI and MDI through project sponsorship. Some of those projects are discussed here, within the context of the physical and chemical properties of TDI and MDI.

It has been known for many years that TDI and MDI can cause respiratory effects at very low concentrations. Accordingly, the production, handling, distribution, use and emission of these materials has been subject to increasingly rigorous control by the industry and regulatory bodies, to protect workers and the population at large. This has given rise to benefits in terms of environmental effects. As a result of engineering controls and well-defined procedures, large spillages are infrequent and usually dealt with effectively, and levels of emission are normally very low.

PRODUCTS AND PROPERTIES

TDI and MDI are supplied to the polyurethane industry as a variety of products, designed to give a range of handling characteristics and polyurethane product properties. These include 80/20-TDI, 65/35-TDI, TDI prepolymers, polymeric MDI, monomeric MDI, and variants of both types of MDI. Of these products 80/20-TDI and polymeric MDI still predominate: some of their physical properties (along with those of monomeric MDI) are given in Table 1.

TDI is sometimes referred to as a "highly reactive and volatile substance." Both points require qualification. The reactivity of TDI (to water and polyols) is normally only observed in catalysed chemical systems used for the production of polyurethanes. We shall see that in the environ-

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Table 1. Some properties of TDI and MDI [1,2,3,4,5].

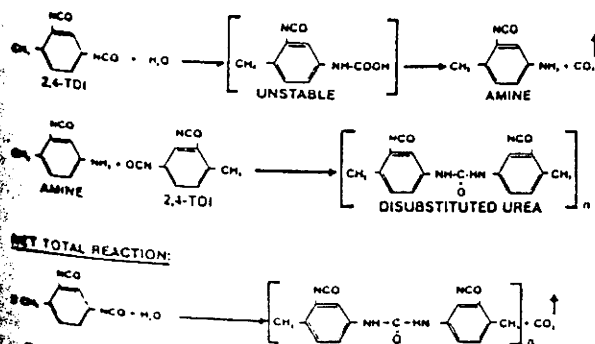
Property		TDI* 80/20	MDI	
			Polymeric	Monomeric
State at 25°C		liquid	liquid	solid
S.G. at 25°C	g/cm ³	1.21	1.23	1.22
Melting Point	°C	ca. 15	< 10	38
Boiling Point	°C	251	Polymerizes at ca 250°C with evolution of CO ₂	171 at 1.33 mbar 200 at 6.6 mbar 230 decomposition
Vapour Pressure				
			(V.P. TDI/ V. P. water)	
	T = 0°C	33 × 10 ⁻⁴	(5 × 10 ⁻⁴)	
	25°C	33 × 10 ⁻³	(10 × 10 ⁻⁴)	< 10 ⁻⁵
	35°C	75 × 10 ⁻³	(13 × 10 ⁻⁴)	< 10 ⁻⁵
Equilibrium Vapour Concentration at 25°C	mg/m ³ (ppm)	220 (30)	ca 0.09 (0.009)	0.09 (0.009)
Flash Point**	°C	135	230	212

*80/20 TDI is 80% 2,4-TDI, 20% 2,6-TDI.

**Cleveland Open Cup, ASTM D92.

ment the rate of reaction of TDI with water depends on a variety of factors. As regards volatility, TDI has much lower equilibrium vapour pressures than does water. Over the range 0–35°C those of TDI are ca 1000 times lower than those of water (see Table 1). At 25°C the equilibrium vapour concentration of TDI is 30 ppm: those for polymeric and pure MDI are considerably lower. The equilibrium vapour concentrations of modified MDIs and TDIs are even lower than those of the parent isocyanates. In Figure 1 is given the generally accepted sequence of reactions following the interaction of TDI with water.

Further reactions will almost certainly take place at the remaining NCO groups. A similar sequence can be illustrated for MDI. The unstable intermediate produced decomposes to the amine with the liberation of CO₂, and the amine reacts immediately with more diisocyanate to yield a polyurea. However, as Saunders and Frisch [6] point out, the interactions of diisocyanates and water are complex and may involve several mechanisms. It is a common misconception that in the presence of water TDI is converted to toluene diamine (TDA) in stoichiometric proportions. This is certainly not the case, but an important question is to what extent TDI (or MDI) gives rise to traces of TDA (or MDA—methylene dianiline) in the environment, in view of the toxic properties of aromatic amines.



EMISSIONS INTO THE AIR

Sources

TDI is used very predominantly for the production of flexible foam slabstock and moulding. Emissions from these processes are known to be richer in 2,6-TDI than is the 80/20 TDI starting material [7]. TDI emissions are often vented to atmosphere, but concentrations are rather low. In a study of six W. German flexible foam factories in 1979, the University of Stuttgart found [8] that stack concentrations were in the range 3–8 mg/m³, which represented about 0.005% of the total TDI used. In the UK and some states of the USA there are very rigorous requirements regarding emissions: "fenceline" concentrations of the order 0.003 mg/m³ (0.0004 ppm TDI) or lower are required in some cases.

As regards MDI, typical emission levels are more difficult to quantify, due to the diversity of applications and wide variety of MDIs (prepolymers and variants) which are used. According to the application the emissions may comprise (a) MDI vapour, (b) MDI aerosol (and vapour), or (c) reacting mix aerosol (and vapour) which will be converted predominantly to a polyurethane. In many applications emission levels are much lower than those from TDI flexible foam processes. About half of the MDI produced is used in moulding (or refrigerator) manufacture, which usually give extremely low emission levels. The British Rigid Urethane Foam Manufacturers' Association has carried out a recent survey [9] of Member Companies' polyurethane production facilities, in which insulation board is produced by spray and liquid laydown techniques, and rigid foam slabstock is produced by both continuous and discontinuous techniques: their production comprises about 50% of total UK rigid foam manufacture. Normal emission levels were found to be 0.2 mg/m³ or lower with occasional excursions above that level.

Developments in polyurethane processing and the control of emissions are leading to improved environmental conditions. Noteworthy here are (a) increasing use of RIM closed-circuit moulding technology and (b) developments in the carbon absorption of emissions [10]. Discussions

[11,12] are in progress in the flexible foam industry to assess the viability of co-absorption of TDI and chlorofluorocarbon emissions, with subsequent recovery of the latter.

The Fate of TDI in the Atmosphere

Several workers [13-16] have carried out studies to investigate the kinetics and reaction products of TDI in the atmosphere. Most of these have been reviewed by Holdren et al. [17]. The results of work in this field should be considered in the light of (a) the highly adsorptive properties of TDI and (b) the possible conversion of TDI to TDA under the conditions of sampling and analysis: similar considerations apply to MDI. Walker and Pinches [18] sampled ambient air in flexible foam factories and concluded that appreciable quantities of TDA had been formed from TDI in the atmosphere. Sandridge [19], in a critique of the study, explained their findings in terms of interfering species in the analyses. Walker acknowledged [20] this possibility and agreed that their conclusions might have been erroneous, or at least, premature. Similar results have not been reported since.

A major study [17,21] on this topic has been carried out by Holdren, Spicer, and Riggan of the Battelle Institute, Columbus, Ohio, U.S.A. Experiments were carried out in a large (17 m³) chamber, lined with PTFE sheeting, in order to minimize wall effects. A series of atmospheres were generated in the chamber to simulate environmental conditions and to determine the parameters giving rise to loss of TDI from the gas phase. Experiments were carried out both in darkness and with irradiation. An important feature of the work was the use of many instrumental techniques to analyse the atmospheres. An assessment of the effects of the following was made (a) photolytic decomposition, (b) photochemically induced pollutants (eg., O₃, OH radicals), (c) urban hydrocarbon mixture and ammonium sulphate particles, (d) TEDA (triethylene diamine), a very commonly used catalyst and (e) possible conversion of TDI to TDA. Outline results of the study are given in Table 2: the final column gives the net loss rates, obtained by subtracting the wall loss rates from the average removal rates.

It was found that under the experimental conditions:

- The first order loss rate of TDI from the vapour phase in humid air (7-70% R.H.) and darkness was rather low (ca. 15% per hour).
- Irradiation caused an increase in loss rate (by ca. 20% per hour), the increase being mainly attributable to free radical attack. The loss rate was little affected by the presence of a variety of common atmospheric pollutants.
- The rate of TDI loss increased very considerably (by 44% per hour) when the level of TEDA vapour was increased from 0.2 ppm to 2 ppm under irradiation conditions.
- No TDA was found above the detection limit of 10 ng/ml, which would correspond to a maximum conversion of 0.05% TDI to TDA.
- Surface absorption onto the chamber lining was a significant removal mechanism.

The above findings indicate that TDI which is emitted during daylight hours has a half-life of about 3 hours, which is little affected by common atmospheric pollutants, and which is independent of relative humidity (7-70%). The loss rate may be affected by the presence of TEDA under factory conditions, although TEDA emission levels are normally well below 2 ppm in flexible foam manufacture, it is believed. There are other tertiary aliphatic amine catalysts, more volatile than TEDA (notably N-ethyl morpholine), which might affect TDI loss rates in practice. A study of emission levels of a range of amine catalysts used in flexible foam technology is currently in progress [22].

Fate of TDA, MDA and TDI under Photolytic Conditions

Theoretical considerations [23] indicate that direct formation of TDA (or MDA) from the corresponding diisocyanates by atmospheric hydrolysis processes is very unlikely, and the Battelle study results support this. Whilst it seemed unlikely that appreciable concentrations of TDA (or MDA) would arise from airborne TDI (or MDI),

Table 2. TDI removal rates.

Experiment	Urban Mix	Irradiation	TEDA	Other Species	Avg. TDI Removal Rate hr ⁻¹	Net Loss Rate (TDI Removal Rate Minus Wall Loss Rate) hr ⁻¹
1	No	No	No	—	0.15*	0
2	No	Yes	No	—	0.36	0.21
3	Yes	Yes	No	—	0.36	0.21
4	Yes	Yes	No	0.5 ppm Ammonia	0.33	0.18
5	Yes	Yes	2 ppm	—	0.99	0.84
6	Yes	Yes	No	100 µg/m ³ Ammonium Sulphate	0.40	0.25
7	No	No	No	—	0.35	0
8	No	Yes	No	4 ppm Nitrous Oxide	0.38	0.03
9	Yes	No	0.2 ppm	—	0.36	0.01
10	Yes	Yes	0.2 ppm	—	0.55	0.20

*0.15/hr = 15%/hr (see text).

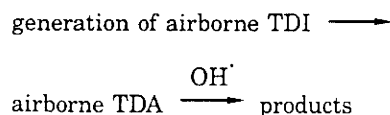
III funded a study to investigate the fate of airborne TDA and MDA, to address their possible formation from any source. The gas phase decomposition of TDI was also investigated. Present knowledge [24] indicates that tropospheric degradation of trace gases (excluding olefinic substances) are predominantly determined by their reactions with OH radicals. (The Battelle study had already indicated that free radical attack is a much more important mechanism than direct photolysis in gas-phase TDI loss.) Accordingly, the study [23], which was carried out by Becker, Bastian and Klein of Wuppertal University, F.R.G., was of OH radical attack. The reaction vessel was a 420 litre glass cylinder into which was introduced the given test substance at atmospheric pressure. Hydroxy radicals were generated by the photolysis of methyl nitrite in the presence of NO to prevent the formation of O₃ and NO₂ radicals. The loss rate of the test substance was compared with that of a reference material at 25°C, using long-path FTIR absorption spectroscopy. The conditions of the experiments were such that the results relate only to gas phase losses, and not to deposition rates or heterogeneous reactions. Decomposition products were not investigated.

Tropospheric half lives (τ) under simulated conditions for the first order bimolecular reaction of the test substance with OH radical (concentration [OH]) were derived from the rate constants k_{OH} , where:

$$\tau = 0.69 (k_{OH} \times [OH])^{-1}$$

The results, along with those of some other substances as cited by Becker and co-workers, are given in Table 3.

The results indicate that under simulated atmospheric conditions the OH radical-initiated reactions of MDA and TDA are relatively fast and more rapid than those of TDI and of several hydrocarbons, for example. Under such conditions, the rate-determining step of a possible sequence:



would be the generation of airborne TDI, and no accumulation of atmospheric TDA would result.

The investigators also studied gas-phase OH radical attack of TDI. The decay rate (0.053 hr⁻¹) was lower than that found by the Battelle Group (0.21 hr⁻¹), but they concluded that this was not unexpected in view of the scatter of results and not fully comparable experimental design. Experiments were carried out at 25°C and 28°C, respectively.

Combustion of TDI and MDI

Fire parameters of TDI and MDI have been studied under laboratory conditions [3,25]. The results are in accordance with practical experience, notably that they are ignited only with difficulty and do not support combustion easily. Their resistance to ignition is reflected in their flash points (Table 1), which are relatively high, compared to those of many products which are transported and stored under similar conditions [3]. Apart from the carefully controlled destruction of TDI and MDI in incinerators, it is likely that they would only be burnt in large acci-

Table 3. Hydroxyl radical attack of various substances.

Substance	Tropospheric 1/2 Lifetimes (hr)
TDI (80:20)	13.0
TDA (2.4-)	0.5
TDA (2.6-)	1.0
MDA	3.2
Propane	82.0
Toluene	15.6
Aniline	0.8

dental fires. It is expected that the combustion products would not be dissimilar to those from a range of natural and synthetic nitrogen-containing compounds, and that no unique harmful products would be formed.

SOIL AND WATER

TDI and MDI may come into contact with soil or water following accidental spillage. Experience gained from such spillages indicates that they are usually well contained. Monomeric MDI (mp 38°C), when handled as a liquid, solidifies on contact with soil or water. Under many circumstances TDI (mp ca. 15°C) and many modified TDIs and MDIs solidify, too. Polymeric MDI solidifies only at low temperatures not usually encountered in the environment. However, polymeric MDI, as well as the other materials under consideration, has specific gravity and viscosity greater than those of water, and experience indicates that it rapidly sinks in water without becoming finely divided. This effect has even been observed in a fast-flowing stream.

Agglomerations of MDI and TDI react with water to form a hard crust of inert, water-insoluble material comprising polyureas. Analysis of such polymeric materials is very difficult and precise work on their composition has not been carried out. However, the products of reaction of polymeric MDI and 80/20 TDI with water investigated in animal studies have been found to give no observable acute effects. LD 50 values for both polymeric MDI- and 80/20 TDI-based polyureas were found [26] to be > 15 g/kg in rats (single gavages in peanut oil, period of observation 14 days, no fatalities).

Soil

Information on the interaction of isocyanates with soil or sand is important in terms of (a) the impact of accidental spillage onto soil and (b) the efficacy and possible environmental effects of using wet soil or sand as a means of decontaminating a spillage area. Large accidental spillages are usually decontaminated by the application of large quantities of water or by covering and mixing the diisocyanate with wet earth. The use of wet earth or sand is preferable, wherever local conditions allow it, because the diisocyanate remains localised and the mixture, when inactive, can be disposed of easily. Washing away material, especially from an impervious surface such as a factory floor or road, could cause further distribution of reacting

Table 4. Analysis of TDI (+ TDA) in soil samples.

After 1 week	TDI (+ TDA) = 0.20 to 100 ppm by wt.
After 6 weeks	TDI (+ TDA) = 0.06 to 1.0 ppm by wt.
After 1 year	TDA not detected (detection limit 0.1 ppm)
After 6 years	TDA not detected at 20-100 cm depth (detection limit 0.05 ppm)

material, and in a more finely divided state if high pressure hosing is used.

Studies on models have been carried out (a) to simulate the covering of a TDI spillage with wet sand and (b) to assess the chemical stability of polyureas prepared from ¹⁴C-labelled MDI and TDI in different agricultural soils. In addition, a study has been carried out on the environmental impact of an actual large spillage of TDI. These have been reviewed elsewhere [27,28], but the main points are outlined below.

The results [29] of model experiments indicated that TDI in undisturbed wet sand (coarse or fine) is converted to polyureas at a rapidly decreasing rate. After 24 hours, 5.5% of the original TDI was unreacted and after 8 days 3.5% remained. These findings can be explained in terms of the encapsulation of TDI within a forming crust of polyurea, which impedes the further penetration of water. No TDA was found above the detection limit of 0.01 ppm. In a separate study [30] the possible degradation of polyureas prepared from ¹⁴C-labelled MDI and TDI was studied in different agricultural soils. No degradation was detected: ¹⁴CO₂ was not evolved, indicating that TDA was not formed.

In April 1975 a road accident occurred, as a result of which 14 tons of TDI were deposited on marshy ground. The spillage was covered with absorbent materials (mainly sand). A six-year study [31] was carried out in close collaboration with the local authority to investigate the consequences of the incident. Outline findings are given in Table 4. No TDI (or TDA) were found in a brook connected to the marsh after intervals of 10 days and 12 weeks.

Analysis of samples at the 1-week and 6-week stages was carried out by a method which did not distinguish between TDI and TDA. It is assumed from the studies reported above that TDI was the predominant species. The results are again compatible with the encapsulation of TDI by a

polyurea crust. It is noteworthy that the vegetation at the site of the incident had developed normally during July 1975: grass grew normally, new foliage appeared on trees and flowers bloomed.

Water

III has funded a number of studies on the chemical and biological effects of MDI, MDA, TDI, and TDA in marine and river water models [32-35]. In addition, Curtis et al. [36] have investigated the toxicity of TDI to freshwater and saltwater organisms. Duff [27], and Brochhagen and Grievesson [28] have reviewed the above findings. Fujiwara [32] carried out studies on the presence of TDI, TDA, MDI, and MDA in marine and river water and in polyurea crusts, following the addition of the respective diisocyanates to the systems. Observations on the river model were made during spring, summer, autumn, and winter. Low concentrations of both diisocyanates and the respective diamines were found in most cases on day 1, but these were transient. It is not foreseen that aquatic life would be subject to long-term exposure from TDI, TDA, MDI, or MDA following a spillage of MDI or TDI. III studies devoted only to the effects of MDA and TDA on aquatic life will be reviewed in a future publication.

Caspers, Hamburger, Kanne, and Klebert [34] of Bayer AG, Leverkusen, F.R.G., have recently completed a comprehensive study for III of the effects of TDI and MDI (also TDA and MDA) on aquatic life, following OECD Guidelines 302C, 209, 202, and 203. Their outline results are presented in Table 5.

The results should be taken as indicators of the general overview of the immediate effects of acute exposure. Details of the methodology and analysis of the results, as presented in the original report, are required for an in-depth interpretation of the study. The findings, which are broadly in agreement with those of other workers [32,33], indicate that:

- The reaction products of TDI and MDI with water do not biodegrade readily.
- TDI and MDI are not appreciably toxic to bacteria.
- When dispersed in water with moderate efficiency, MDI and TDI are not appreciably toxic to daphnia: no negative effects on their reproduction were found at the highest concentrations used.
- Results on the toxicity of MDI and TDI to fish were rather inconsistent and the authors comment that harmful effects due to oral ingestion or mechanical violation of body tissues could not be excluded. The broad finding was that the immediate toxic effects of MDI and TDI due to acute exposure are rather low.

The investigators carried out several tests with very high shear stirring, and found increased fish and daphnia toxicity under such conditions: the results are not included here since such conditions could not be foreseen in the environment. A simple understanding of the acute fish toxicity of TDI and MDI can not be gained from the different LC 50 results of Hamburger et al., obtained at different stirring rates, taken along with the results of Fujiwara and Curtis et al., who each used different species and different experimental conditions. This is not unexpected. Such studies with MDI and TDI are especially difficult to interpret because of the inherent problem that the chemicals are almost totally insoluble in, and react with, the

Table 5. Results of Hamburger and co-workers.

Test	MDI	TDI
Biodegradation (Inherent) % in 28 days	None	None
Bacteria Toxicity (E. Coli) mg/l, 24 h	EC 50 > 100	EC 50 > 100
Daphnia Reproduction (Daphnia magna) mg/l, 24 h	no negative effects at highest concentration (TDI - 0.5, MDI - 10)	
Daphnia Toxicity (Daphnia magna) mg/l, 24 h	EC 50 ≥ 1000	EC 50 ~ 750
Fish Toxicity (Zebra fish) mg/l, 96 h	LC 0 > 1000	LC 0 > 100 LC 100 ≥ 250

medium to form insoluble products. OECD Guidelines do not define the mode of addition of such materials. Under different stirring conditions the physical form and the chemical composition of the reaction products will differ. In that respect it is interesting to note the comment of Curtis and co-workers [36] who found TDI hazardous to freshwater minnows (but not to saltwater shrimp): "The TDI appeared to be toxic to fathead minnows only in unreacted form, since all mortalities occurred during the first twelve hours of test. A concurrent decrease in pH was observed as a result of carbon dioxide formation." It is also possible that toxic effects could have resulted from the associated formation of carbon dioxide.

CONCLUSIONS

This paper suggests that the overall level of environmental pollution from TDI and MDI is very low. Emission levels are low and spillages of MDI or TDI are usually localized, and the diisocyanates very largely converted to materials which are chemically and biologically inert. There is, however, scope for further reductions of emissions or spillages, especially by those users who do not observe rigorous procedures for handling TDI and MDI.

When viewed in their entirety the investigations cited above provide an insight into the probable effects of MDI and TDI in the environment. The evidence indicates that the ecological impact is likely to be slight, and reversible. However, it is recognized that there are limitations to the reported studies of environmental effects. There are many difficulties inherent in the extrapolation from model systems to actual cases; also there are limitations to analytical techniques. Accordingly, the ICI continues its researches in this field in the interests of man and the environment.

ACKNOWLEDGEMENT

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BIOGRAPHY

David S. Gilbert



and now works exclusively for the III.

Having graduated in industrial chemistry, David Gilbert undertook research studies of the kinetics of organic chemical reactions, notably using radiochemical techniques to follow symmetrical exchange reactions. Most of his career has been in polyurethanes, working with ICI on elastomers, flexible foams and rigid foams. In 1982 he established the consultancy David Gilbert Associates

*A. Chinn
O. Hale
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Dow Book*

MATERIAL SAFETY DATA SHEET

Dow Chemical U.S.A.* Midland, MI 48674 Emergency Phone: 517-636-4400

Product Code: 92097

Page: 1

PRODUCT NAME: VORANATE (R) T-80 TYPE I TOLUENE DIISOCYANATE

Effective Date: 12/13/88 Date Printed: 01/12/89

MSDS:000609

1. INGREDIENTS: (% w/w, unless otherwise noted)

Toluene-2,4-diisocyanate (TDI)	CAS# 000584-84-9	80%
Toluene-2,6-diisocyanate	CAS# 000091-08-7	20%

This document is prepared pursuant to the OSHA Hazard Communication Standard (29 CFR 1910.1200). In addition, other substances not 'Hazardous' per this OSHA Standard may be listed. Where proprietary ingredient shows, the identity may be made available as provided in this standard.

2. PHYSICAL DATA:

BOILING POINT: 250C (482F)
VAP PRESS: 0.01 mmHg @ 20C
VAP DENSITY: 6.0
SOL. IN WATER: Insoluble
SP. GRAVITY: 1.22 @ 25/15.5C
APPEARANCE: Water white to pale yellow liquid.
ODOR: Sharp pungent odor.

3. FIRE AND EXPLOSION HAZARD DATA:

FLASH POINT: 127C (260F)
METHOD USED: PMCC, ASTM D-93

FLAMMABLE LIMITS

LFL: Not determined
UFL: Not determined

EXTINGUISHING MEDIA: Carbon dioxide, dry chemical, or foam.
If water is used, it should be in very large quantity.
The reaction between water and hot isocyanate may be vigorous.

FIRE & EXPLOSION HAZARDS: Down-wind personnel must be evacuated.

(Continued on Page 2)

(R) Indicates a Trademark of The Dow Chemical Company

M A T E R I A L S A F E T Y D A T A S H E E T

Dow Chemical U.S.A.* Midland, MI 48674 Emergency Phone: 517-636-4400

Product Code: 92097

Page: 2

PRODUCT NAME: VORANATE (R) T-80 TYPE I TOLUENE DIISOCYANATE

Effective Date: 12/13/88 Date Printed: 01/12/89

MSDS:000609

3. FIRE AND EXPLOSION HAZARD DATA: (CONTINUED)

Do not reseal contaminated containers since pressure build-up may cause rupture. Fire point: 146C (295F).

FIRE-FIGHTING EQUIPMENT: People who are fighting isocyanate fires must be protected against nitrogen oxide fumes and isocyanate vapors by wearing positive pressure self-contained breathing apparatus and full protective clothing.

4. REACTIVITY DATA:

STABILITY: (CONDITIONS TO AVOID) Stable when stored under recommended storage conditions. Store in a dry place at temperatures between 18-41C (65-105F).

INCOMPATIBILITY: (SPECIFIC MATERIALS TO AVOID) Water, acid, base, alcohols, metal compounds, surface active materials. Avoid water as it reacts to form heat, CO₂ and insoluble urea. The combined effect of the CO₂ and heat can produce enough pressure to rupture a closed container.

HAZARDOUS DECOMPOSITION PRODUCTS: Isocyanate vapor and mist, carbon dioxide, carbon monoxide, nitrogen oxides and traces of hydrogen cyanide.

HAZARDOUS POLYMERIZATION: May occur with incompatible reactants, especially strong bases, water or temperatures over 41C (105F).

5. ENVIRONMENTAL AND DISPOSAL INFORMATION:

ACTION TO TAKE FOR SPILLS/LEAKS:

Evacuate and ventilate spill area, dike spill to prevent entry into water system, wear full protective equipment including respiratory equipment during clean up.

Major spill: Call Dow Chemical U.S.A. (409) 238-2112. If

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M A T E R I A L S A F E T Y D A T A S H E E T

Dow Chemical U.S.A.* Midland, MI 48674 Emergency Phone: 517-636-4400

Product Code: 92097

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PRODUCT NAME: VORANATE (R) T-80 TYPE I TOLUENE DIISOCYANATE

Effective Date: 12/13/88 Date Printed: 01/12/89

MSDS:000609

5. ENVIRONMENTAL AND DISPOSAL INFORMATION: (CONTINUED)

transportation spill involved call CHEMTREC (800) 424-9300. If temporary control of isocyanate vapor is required a blanket of protein foam (available at most fire departments) may be placed over the spill. Large quantities may be pumped into closed but not sealed containers for disposal.

Minor spill: Absorb the isocyanate with sawdust or other absorbent and shovel into open top containers. Do not make pressure tight. Transport to a well-ventilated area (outside) and treat with neutralizing solution consisting of a mixture of water and 3-8% concentrated ammonium hydroxide or 5-10% sodium carbonate. Add about 10 parts of neutralizer per part of isocyanate with mixing. Allow to stand for 48 hours letting evolved carbon dioxide to escape.

Clean-up: Decontaminate floor using water/ammonia solution with 1-2% added detergent letting stand over affected area for at least 10 minutes. Cover mops and brooms used for this with plastic and dispose properly (often by incineration).

DISPOSAL METHOD: Follow all federal, state and local regulations. Liquids are usually incinerated in a proper facility. Solids are usually also incinerated or landfilled. Empty drums should be filled with water. Let drum stand unsealed for 48 hours. Before disposal drums should be drained, triple rinsed, and holed to prevent reuse. Dispose of drain and rinse fluid according to federal, state and local laws and regulations. The most commonly accepted method is in an approved wastewater treatment facility. Drums should be disposed of in accordance with federal, state and local laws and regulations. Commonly accepted methods for disposal of plastic drums are disposal in an approved landfill after shredding or incineration in an approved industrial incinerator or other appropriate incinerator facility. Steel drums are commonly disposed in an approved landfill after crushing or in accordance with other approved procedures.

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M A T E R I A L S A F E T Y D A T A S H E E T

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PRODUCT NAME: VORANATE (R) T-80 TYPE I TOLUENE DIISOCYANATE

Effective Date: 12/13/88 Date Printed: 01/12/89

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6. HEALTH HAZARD DATA:

EYE: May cause pain, severe eye irritation and moderate corneal injury. Vapors may irritate eyes.

SKIN CONTACT: Prolonged or repeated exposure may cause severe irritation, even a burn. Skin contact may result in allergic reaction even though it is not expected to result in absorption of amounts sufficient to cause other adverse effects.

SKIN ABSORPTION: The LD50 for skin absorption in rabbits is >9400 mg/kg.

INGESTION: Single dose oral toxicity is low. The oral LD50 for rats is 5800 mg/kg. Ingestion may cause gastrointestinal irritation or ulceration.

INHALATION: Excessive vapor concentrations are attainable and could be hazardous on single exposure. Single and repeated excessive exposure may cause severe irritation to upper respiratory tract and lungs (choking sensation, chest tightness), respiratory sensitization, decreased ventilatory capacity, liver effects, cholinesterase depression, gastrointestinal distress and/or neurologic disorders. The 4-hour LC50 for TDI for rats is 13.9 ppm.

SYSTEMIC & OTHER EFFECTS: Based on available data, repeated exposures are not anticipated to cause any additional significant adverse effects. For hazard communication purposes under OSHA standard 29 CFR Part 1910.1200, this chemical is listed as a potential carcinogen by Nat'l. Tox. Program and IARC. An oral study in which high doses of TDI were reported to cause cancer in animals has been found to contain numerous deficiencies which compromise the validity of the study. TDI did not cause cancer in laboratory animals exposed by inhalation, the most likely route of exposure. Birth defects are unlikely. Exposures having no effect on the mother should have no effect on the fetus. Did not cause birth defects in animals; other effects were seen in the fetus only at doses which caused toxic effects to the mother. Results of in vitro ("test tube") mutagenicity

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MATERIAL SAFETY DATA SHEET

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PRODUCT NAME: VORANATE (R) T-80 TYPE I TOLUENE DIISOCYANATE

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6. HEALTH HAZARD DATA: (CONTINUED)

tests have been inconclusive.

7. FIRST AID:

EYES: Irrigate with flowing water immediately and continuously for 15 minutes. Consult medical personnel.

SKIN: In case of contact, immediately flush skin with plenty of water for at least 15 minutes while removing contaminated clothing and shoes. Call a physician if irritation persists. Wash clothing before reuse. Destroy contaminated shoes.

INGESTION: Do not induce vomiting. Call a physician and/or transport to emergency facility immediately.

INHALATION: Remove to fresh air. If not breathing, give mouth-to-mouth resuscitation. If breathing is difficult, give oxygen. Call a physician.

NOTE TO PHYSICIAN: May cause tissue destruction leading to stricture. If lavage is performed, suggest endotracheal and/or esophagoscopy control. If burn is present, treat as any thermal burn, after decontamination. No specific antidote. Supportive care. Treatment based on judgment of the physician in response to reactions of the patient. The manifestations of the respiratory symptoms, including pulmonary edema, resulting from acute exposure may be delayed. May cause respiratory sensitization. Cholinesterase inhibition has been noted in human exposure but is not of benefit in determining exposure and is not correlated with signs of exposure.

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M A T E R I A L S A F E T Y D A T A S H E E T

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PRODUCT NAME: VORANATE (R) T-80 TYPE I TOLUENE DIISOCYANATE

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8. HANDLING PRECAUTIONS:

EXPOSURE GUIDELINE(S): OSHA PEL is 0.02 ppm as a ceiling limit for toluene 2,4-diisocyanate. ACGIH TLV is 0.005 ppm; 0.02 ppm STEL for toluene 2,4-diisocyanate. Dow Industrial Hygiene Guide is 0.02 ppm as a ceiling limit for toluene diisocyanate.

VENTILATION: Provide general and/or local exhaust ventilation to control airborne levels below the exposure guidelines.

RESPIRATORY PROTECTION: Atmospheric levels should be maintained below the exposure guideline. When respiratory protection is required for certain operations, use an approved supplied-air respirator. For emergency and other conditions where the exposure guideline may be greatly exceeded, use an approved positive-pressure self-contained breathing apparatus.

SKIN PROTECTION: Use protective clothing impervious to this material. Selection of specific items such as gloves, boots, apron, or full-body suit will depend on operation. Remove contaminated clothing immediately, wash skin area with soap and water, and launder clothing before reuse. Safety shower should be located in immediate work area.

EYE PROTECTION: Use chemical goggles. If vapor exposure causes eye irritation, use a full-face, supplied-air respirator. Eye wash fountain should be located in immediate work area.

9. ADDITIONAL INFORMATION:

REGULATORY REQUIREMENTS:

SARA HAZARD CATEGORY: This product has been reviewed according to the EPA 'Hazard Categories' promulgated under Sections 311 and 312 of the Superfund Amendment and Reauthorization Act of 1986 (SARA Title III) and is considered, under applicable definitions, to meet the following categories:

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M A T E R I A L S A F E T Y D A T A S H E E T

Dow Chemical U.S.A.* Midland, MI 48674 Emergency Phone: 517-636-4400

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PRODUCT NAME: VORANATE (R) T-80 TYPE I TOLUENE DIISOCYANATE

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9. ADDITIONAL INFORMATION: (CONTINUED)

An immediate health hazard
A delayed health hazard
A reactive hazard

SPECIAL PRECAUTIONS TO BE TAKEN IN HANDLING AND STORAGE: Warning properties of this material (irritation of eyes, nose and throat) not adequate to prevent chronic overexposure from inhalation. This material can produce asthmatic sensitization upon either single inhalation exposure to a relatively high concentration or upon repeated inhalation exposure to lower concentrations. Exposures to vapors of heated TDI can be

extremely dangerous. (Have TDI neutralizer available for spills.)

MSDS STATUS: Revised Section 9

SARA 313 INFORMATION:

This product contains the following substances subject to the reporting requirements of section 313 of Title III of the Superfund Amendments and Reauthorization Act of 1986 and 40 CFR Part 372:

CHEMICAL NAME	CAS NUMBER	CONCENTRATION
TOLUENE-2,6-DIISOCYANATE	000091-08-7	20 %
TOLUENE-2,4-DIISOCYANATE	000584-84-9	80 %

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The Information Herein Is Given In Good Faith, But No Warranty, Express Or Implied, Is Made. Consult The Dow Chemical Company For Further Information.

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